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TMI-2 CORE DEBRIS GRAB SAMPLES --**EXAMINATION AND ANALYSIS** PART 1

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Prepared for the

U.S. Department of Energy Three Mile Island Operations Office Under Contract No. D€-AC07-76ID01570

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TMI-2 CORE DEBRIS GRAB SAMPLES--EXAMINATION AND ANALYSIS

PART 1

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Published September 1986

EG&G Idaho, Inc. Idaho Falls, Idaho 83415

Prepared for the
U.S. Department of Energy
Idaho Operations Office
Under DOE Contract No. DE-ACO7-76ID01570

ABSTRACT

Six samples of particulate debris were removed from the TMI-2 core rubble bed during September and October 1983, and five more samples were obtained in March 1984. The samples (up to 174 g each) were obtained at two locations in the core: H8 (center) and E9 (mid-radius). Ten of the eleven samples were examined at the Idaho National Engineering Laboratory to obtain data on the physical and chemical nature of the debris and the postaccident condition of the core. Portions of the samples also were subjected to differential thermal analysis at Rockwell Hanford Operations and metallurgical and chemical examinations at Argonne National Laboratories. This report presents results of the examination of the core debris grab samples, including physical, metallurgical, chemical, and radiochemical analyses. The results indicate that temperatures in the core reached at least 3100 K during the TMI-2 accident, fuel melting and significant mixing of core structural material occurred, and large fractions of some radionuclides (e.g., 90Sr and 144Ce) were retained in the core.

SUMMARY

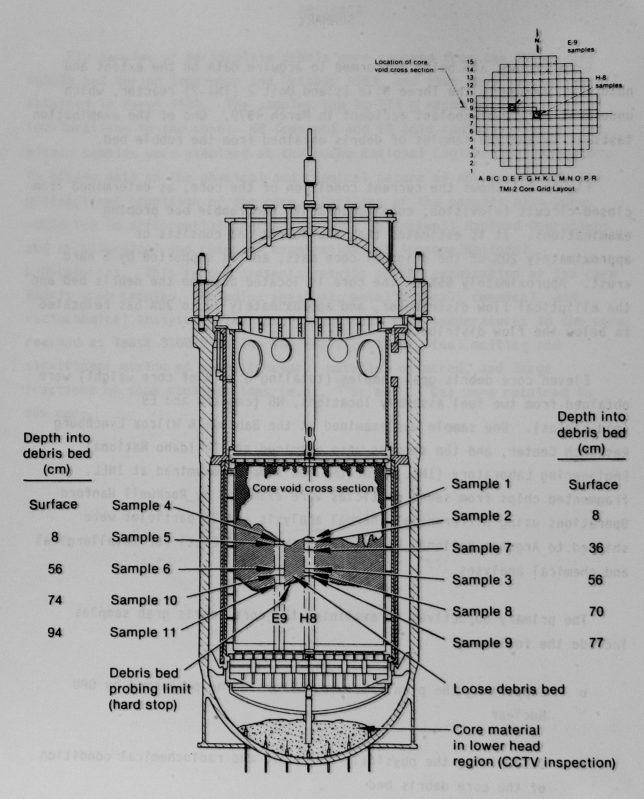
Examinations are being performed to acquire data on the extent and nature of damage to the Three Mile Island Unit 2 (TMI-2) reactor, which underwent a loss-of-coolant accident in March 1979. One of the examination tasks is to analyze samples of debris obtained from the rubble bed.

figure S-1 shows the current condition of the core, as determined from closed-circuit television, core topography, and rubble bed probing examinations. It is estimated that the debris bed consists of approximately 20% of the original core mass, and is supported by a hard crust. Approximately 65% of the core is located between the debris bed and the elliptical flow distributor, and approximately 10 to 20% has relocated to below the flow distributor.

Eleven core debris grab samples (totaling 0.001% of core weight) were obtained from two fuel assembly locations, H8 (center) and E9 (mid-radius). One sample was examined at the Babcock & Wilcox Lynchburg Research Center, and ten samples were examined at the Idaho National Engineering Laboratory (INEL). Of the ten samples examined at INEL, fragmented chips from seven particles were examined by Rockwell Hanford Operations using differential thermal analysis and 22 particles were shipped to Argonne National Laboratories (West and East) for metallurgical and chemical analyses.

The primary objectives of examining the core debris grab samples include the following:

- o Supporting the plant recovery effort being performed by GPU Nuclear
- o Determining the physical, chemical, and radiochemical condition of the core debris bed



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Figure S-1. Schematic showing the current known condition of the TMI-2 core and the locations of the core debris grab samples.

Providing data to support assessment of severe fuel damage codes, primarily with regard to core damage progression and fission product behavior.

This report presents the results and conclusions of analysis of the core debris grab samples, including: (a) the sample acquisition, examination plan, and analytical methods; (b) examination results (physical, metallurgical, chemical, and radiochemical); (c) an evaluation of the overall core condition and radionuclide behavior, using information from the core debris and other available examination results; and (d) principal observations and conclusions obtained from these examinations.

Sample Acquisition, Examination Plan, and Analytical Methods

The core debris grab samples were removed from the debris bed at depths ranging from the surface to 77 cm at the H8 location, and from the surface to 94 cm at the E9 location. A hard stop was encountered at 77 cm at the H8 location, and friction prevented penetration of the debris bed beyond 94 cm at E9. Sample weights obtained ranged from 17 g (Sample 4) to 174 g (Sample 10).

figure S-2 is a flow diagram showing the examinations performed on the core debris grab samples. There were four general types of samples examined: (a) bulk samples, as received from TMI-2; (b) particle size fractions—subgroupings of the bulk samples by size, obtained by sieving the bulk samples, (c) recombined bulk samples—approximately one—third of each particle size fraction, recombined to approximate the original bulk sample; and (d) particles and aliquots—individual particles from the larger (>1000 µm) particle size fractions and aliquots [representative portions from the smaller (<1000 µm) particle size fractions]. Analytical methods used for the examinations are standard examination techniques, with some modifications necessitated by the high radiation fields and insoluble nature of the core debris.

Figure S-2. Flow diagram showing the examinations performed on the TMI-2 core debris grab samples (typical for each sample).

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Physical Examinations

The physical examinations included visual and photographic examinations, weighing, bulk tap density measurements, particle size distribution analysis, and ferromagnetic and pyrophoricity tests. In general, the core debris was composed of particulate material that was not identifiable as core components. Five categories of particles were identified based on visual appearance: (a) apparent fuel pieces; (b) cladding chunks, (c) foamy/porous, prior molten material; (d) particles that are a composite of fuel and prior molten material; and (e) metallic appearing, prior molten particles.

Bulk tap density measurements indicate that there are two density ranges for the samples. Samples 1, 3, and 6 have densities ranging from 3.5 to 3.8 g/cm^3 whereas Samples 9. 10. and 11 have densities ranging from 5.0 to 5.5 g/cm^3 . The particle size distribution analysis indicated that Samples 1 and 3 have the largest fraction of material in the 1680 to 4000-um particle size range, whereas Samples 9, 10, and 11 show a bimodal distribution, with peaks at the 1680 to 4000-um and 297 to 707-um sizes. The bi-modal distribution probably resulted in more efficient packing of the debris material and may explain the higher densities of Samples 9. 10, and 11. The particle size distribution analysis indicates the presence of mostly (>80%) larger particles (>1000 µm). Stratification of the debris bed into two layers also is indicated by the particle size distribution: a surface layer, and another beginning between 36 cm and 56 cm below the debris bed surface. The lower layer contains larger quantities of smaller sized particles. The measurable ferromagnetic material content of Sample 6, the only sample analyzed for ferromagnetic content, was 0.9% of the total sample weight.

Metallurgical Examinations

Twenty-nine particles were selected from the ten core debris grab samples for detailed metallurgical analysis, which included optical and scanning electron microscopy for microstructural information and energy

dispersive x-ray spectroscopy and scanning Auger spectroscopy for information on elemental composition.

Most of the particles examined contained regions of prior molten U-Zr-O, indicating peak temperatures of greater than 2200 K. Of these, many are prior molten $(U,Zr)O_2$, indicating peak temperatures greater than 2800 K. There are a few examples of prior molten material that are almost pure UO_2 , indicating temperatures up to 3100 K (the effect of small amounts of impurities on the melting point of UO_2 is not well known). However, much of the fuel probably remained at fairly low temperatures (<2000 K) relative to the peak temperature or was exposed to high temperatures for only a short time. This conclusion is based on the relatively unrestructured appearance observed for much of the fuel, even though there are attached regions of prior molten ceramics.

The regions of prior molten U-Zr-O usually contain at least some traces of non-fuel rod materials (Al, Cr, Fe, and Ni). These elements often were observed at grain boundaries or in voids. For some particles, they are a significant portion of the total particle weight. Control rod materials (Ag, In, and Cd) were not as commonly found as structural materials. However, major portions of three particles are metallic Ag containing Ni-Sn inclusions, indicating that control rod materials (or at least Ag) apparently had less of a tendency to mix or interact with fuel rod materials than did the structural materials.

<u>Chemical Examinations</u>

Elemental analysis of the core debris grab samples was performed by inductively coupled plasma spectroscopy (ICP) on the recombined bulk samples and the individual particles and aliquots. Elements for which analyses were performed were selected to characterize five groups of core components: (a) uranium fuel and zircaloy cladding, (b) Ag-In-Cd control rod materials, (c) poison rod materials (B, Gd, and Al), (d) structural materials (stainless steel and Inconel) and (e) elemental Te.

Most elemental constituents of the core components were present in all recombined bulk samples, particles, and aliquots, suggesting that significant disruption, mixing, and relocation of these components has occurred. The analyses for U and Zr indicate that significant depletion ($\leq 50\%$) of Zr from the debris bed has occurred. Analyses for control rod materials with emphasis on Ag indicate that as much as 90% of the Ag has relocated from the debris bed. There appears to be a general increase in Ag concentration associated with a decrease in particle size. The burnable poison rod materials, Al and Gd, were measured in most of the samples. The wide distribution of Gd is of particular interest, because only four fuel assemblies in the core contained Gd control material (13 kg). These fuel assemblies were located at four quadrants, about mid-radius in the core.

All is concentrated at the surface of the debris bed, suggesting that the debris bed may have functioned as a trap or retention zone for some core materials. Structural materials are well mixed in the debris bed in concentrations similar to their original concentrations in the fueled portion of a fuel assembly. Elemental Te, which may originate from natural or fission product sources is concentrated near the surface of the debris bed similar to aluminum.

Radiochemical Examinations

Radiochemical examinations were performed on the recombined bulk samples, particles and aliquots from all samples, and the ferromagnetic components of Sample 6. The examinations included gamma spectroscopy, fissile/fertile material (235 U/ 238 U) content, and, 129 I and 90 Sr analyses. Examinations were performed to characterize radionuclides of expected high volatility (129 I and 137 Cs), intermediate volatility (106 Ru and 125 Sb) and low volatility (90 Sr and 144 Ce).

Comparison of the measured to predicted retention/release of these radionuclides in the core debris is shown in Table S-1 and approximately follows the expected volatility of the radionuclides (i.e., the most volatile materials, $\frac{129}{1}$, and $\frac{137}{1}$ Cs, were most significantly

TABLE S-1. RADIONUCLIDE RETENTION NORMALIZED TO URANIUM CONTENT (% of core inventory)

	Calculated Core Average Concentration ^a (C1/g)	H8 Sam	ples	E9 Samples		
<u>Radionuclide</u>		Measured Average <u>Retention^b</u>	_Range ^C	Measured Average Retention ^b	Range ^C	
90 _{Sr}	8.12E-3	94	91-102	92	79-100	
106 _{Ru}	1.41E-3	49	35-74	63	52-86	
125 _{Sb}	4.53E-4	28	19-37	28	18-38	
129 _I	2.81E-9	24 ^d	19-28	19 ^d	10-25	
137 _{Cs}	9.32E-3	19	18-21	24	6-32	
144 _{Ce}	3.37E-3	121	114-128	107	90-130	
235 _{U/U} e		2.5	2.3-2.7	2.7	2.5-2.8	

a. Calculated based on ORIGEN-2 analyses, decays corrected to April 1, 1984.

b. Average measured retention for all samples.

c. Range of retention for recombined bulk samples.

d. Calculated based on particle and aliquot analyses. Uncertainty is $\sim 100\%$.

e. Uranium enrichment calculated from fissile and total U analyses.

released). The results shown are normalized to predicted core average concentrations and are useable for intercomparison purposes. Actual incore fission product concentrations per gram of fuel may vary by a factor of 2. However, the retentions of the low volatile fission products (e.g., $^{90}\mathrm{Sr}$ and $^{144}\mathrm{Ce}$) near 100% suggest that the listed retentions are accurate within 20 to 30%. The high concentrations of $^{144}\mathrm{Ce}$ are probably due to the presence of localized zones of higher burnup fuel at the locations sampled. The $^{129}\mathrm{I}$ content of the samples tends to increase with surface area for the small particle sizes and may indicate a surface area retention dependent mechanism. Also, a correlation was observed between the concentrations of N1 and the $^{125}\mathrm{Sb}$ and $^{106}\mathrm{Ru}$ concentrations, suggesting that N1 functioned as a scavenging material for these radionuclides in the debris.

Contributions to Understanding the TMI Accident

This section describes how the results from the examination of the core debris grab samples contribute to understanding the condition of the TMI-2 core and accident scenario.

Metallurgical analysis of the large particles from the core debris grab samples indicate in some of the material that peak temperatures up to the melting point of the UO₂ fuel were achieved. However, a significant amount of material showed no restructuring, indicating temperatures below about 2200 K. There are indications of either a prolonged candling sequence or multiple temperature excursions. Evidence of movement of molten material down the fuel-cladding gap was observed, and there is some evidence of strong interaction of the fuel materials.

a. The term "candling" is used to denote melting, relocation, and resolidification, similar to that which occurs to the wax of a lighted candle as it is burned.

Evaluations were performed to estimate the degree of mixing, physical relocation, and segregation of core materials. The examination results suggest that the molten core debris was mixed vigorously to produce the degree of mixing and homogenity observed. However, the observed mixing may be localized phenomea at the locations sampled and may not be representative of the entire debris bed.

Evidence of physical relocation and mixing of the fuel assembly materials was obtained by comparing the original U enrichments of the H8 and E9 fuel assemblies with the measured enrichments of samples removed from these locations. No correlation exists between the original and measured enrichments, indicating significant physical relocation of fuel at the H8 and E9 locations occurred.

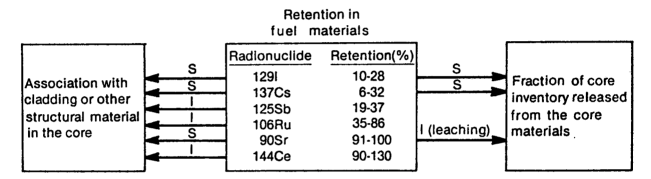
There is evidence of significant relocation and segregation of some non-fuel core materials (Zr, Ag, In, Cd, Gd, and Al). The contributions of these material movements on understanding the core condition and accident scenario are:

- o The unexamined portion of the core may have zones of heterogeneous material composition with significantly different compositions than the debris bed samples.
- o Accumulations of Al and Te at the surface of the debris bed suggest that the surface of the debris bed functioned as a trap for some structural materials. Several mechanisms have been suggested that would allow the observed segregation to have occurred (a) during the accident (vaporization and subsequent condensation on surfaces) or (b) during natural recirculation after the accident (deposition of AlO (OH) or Al₂O₃ powder on the surface of the debris bed).
- The concentrations of Ag tend to be higher for the smaller (<1000 μ m) particle size fractions, suggesting either finely fragmented Ag or a retention mechanism that is surface area dependent.

 90 Sr was retained to a greater extent and 106 Ru to a lesser extent than was expected, based on the volatility of the elements. The greater 90 Sr retention in the samples may be attributable to the formation of SrO, a highly involatile oxide of Sr. The greater release of 106 Ru may be attributed to the formation of highly volatile and quite unstable oxides, RuO₂ and RuO₄.

Some radionuclides exhibited significant mobility from the fuel to other core materials and to the remainder of the reactor coolant system. Significant fractions of the core inventories of 137 Cs and 129 I have been measured outside the reactor core. Transport or retention mechanisms for 125 Sb, 90 Sr, and 106 Ru are discussed above. The transport of 144 Ce from the fuel to structural materials is important, because this radionuclide is expected to be strongly retained in the fuel. A suggested mechanism for the possible transport of this radionuclide from the fuel is the formation of CeO, a relatively volatile oxide of Ce. However, formation of this compound requires that $\mathrm{Ce_2O_3}$ is reduced and would suggest that periods of oxidizing and reducing atmospheres occurred during the accident. Reducing atmospheres may have occurred during periods of no coolant or in isolated portions of the core during secondary temperature ramps probably after 150 minutes into the accident.

The data obtained from analysis of the core debris grab samples suggest that the condition of the lower regions of the core might be significantly different than that observed in the debris bed. The data also suggest that radionuclides might be present as accumulations or that there may be depleted regions lower in the core.



S - Significant relocation
I - Insignificant relocation
5 3266

Figure S-3. Schematic showing the behavior (retention/relocation) of various radionuclides in the TMI-2 core.

ACKNOWLEDGMENTS

Many people contributed to the analysis of the core debris grab samples. Special thanks go to the following individuals: Roger H. Wilhelmsen and Alan W. Marley for assisting in the analysis work: Robert J. Gehrke and the staff at the Radiation Measurement Laboratory for their timely gamma spectroscopy analysis of the samples: Yale D. Harker and Paul R. Napper for developing the fast neutron flux measurement facility and performing the fissile/fertile material analysis: Joseph E. Stoyack and Michael Benton for their assistance in conducting the metallography: Daniel A. Pavlica and Theodore R. Lyon of Westinghouse Idaho Nuclear Company, Inc. for performing inductively coupled plasma, Auger, and energy dispersive x-ray spectroscopy measurements and scanning electron microscopy (SEM) examinations on particles from the first group of grab samples; Bertil G. Carlson of Argonne National Laboratory-West for performing optical metallography and John Sanecki and Henry Hoff of Argonne National Laboratory-East (ANL-E) for performing SEM/EDS and SAS examinations on 22 particles from the two groups of grab samples; Greg R. Eldam of GPU Nuclear for his contribution to the particle size evaluations; Dr. Peter Hofmann of KfK in the Federal Republic of Germany, Bob Strain and Larry Neimark of ANL-E, George Hayner of B&W, and M. L. Picklesimer for their invaluable, indepth review and analysis of the metallography results; Richard R. Hobbins for his assistance in analyzing the data and reviewing the manuscript; Yoshiaki Izumoji for Appendix G; Takeshi Yamahara for technical review: and Angela K. Richardson for assistance in organizing Appendix C.

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NOME NCLATURE

Allquot A representative portion of a homogeneous

particle size fraction.

Bulk tap density Density of the bulk sample obtained by placing

the sample in a graduated beaker, tapping the beaker to compact the material, measuring the volume, and dividing the sample weight (g) by

the volume (cm^3) .

Fuel bundle Fuel assembly without end fittings.

Particle A piece of debris. Analysis was performed on

individual particles greater than 1000 μm in size. For particles smaller than 1000 μm ,

analysis was performed on aliquots.

Particle size fractions Subgroupings of the samples by particle size,

resulting from the sleving operation. The particle sizes range from less than 20 to

greater than 4000 µm.

Recombined bulk sample After the particle size analysis,

approximately one-third of each particle size

fraction was weighed, then recombined to

approximate the original sample.

Sample All debris contained in a sampling tool.

as-received from GPU Nuclear.

ACRONYMS AND ABBREVIATIONS

BSE Backscatter electron analysis

CCTV Closed-circuit television

EDS Energy dispersive x-ray spectroscopy

ICP Inductively coupled plasma spectroscopy

SAS Scanning Auger spectroscopy

SEM Scanning electron microscopy

Particle 1A Example of notation used to identify the

particles (Particle 1A was the first of several particles selected from Sample 1).

Sample 3 (H8, 56 cm) *

Example of notation used to identify the samples and their original location in the debris bed (Sample 3 was obtained from the H8 core location, 56 cm into the debris bed).

DRAFT REPORT: TMI-2 CORE DEBRIS GRAB SAMPLES EXAMINATION AND ANALYSIS

PART 1

1. INTRODUCTION

The Unit 2 pressurized water reactor at Three Mile Island (TMI-2) underwent a prolonged loss-of-coolant accident on 28 March 1979, that resulted in severe damage to the core. As a consequence of the TMI-2 accident, numerous aspects of light water reactor safety have been questioned, and the U.S. Nuclear Regulatory Commission (NRC) has embarked on a thorough review of reactor safety issues, particularly the causes and effects of severe core damage accidents. The nuclear community acknowledges the importance of examining TMI-2 in order to understand the nature of the core damage. Immediately after the accident, four organizations with interests in both plant recovery and acquisition of accident data formally agreed to cooperate in these areas. These organizations, General Public Utilities Nuclear Corporation (GPU Nuclear--owner/operator of TMI). Electric Power Research Institute (EPRI). the U.S. Nuclear Regulatory Commission (NRC), and the U.S. Department of Energy (DOE), collectively known as GEND, are presently involved in postaccident evaluation of TMI-2. DOE is providing a portion of the funds for reactor recovery (in those areas where accident recovery knowledge will be of generic benefit to the light water reactor industry of the United States). In addition, DOE is funding acquisition and analysis of severe accident data obtained from examining the damaged core.

The present condition of the TMI-2 core, as determined by closed-circuit television (CCTV) examination and probing of the debris bed, is illustrated in Figure 1. Current estimates indicate that approximately 20% of the total core mass is an upper layer of rubblized debris supported by a hard crust. Approximately 65% is located between the debris bed and the elliptical flow distributor, and approximately 10 to 20% has relocated to below the flow distributor.

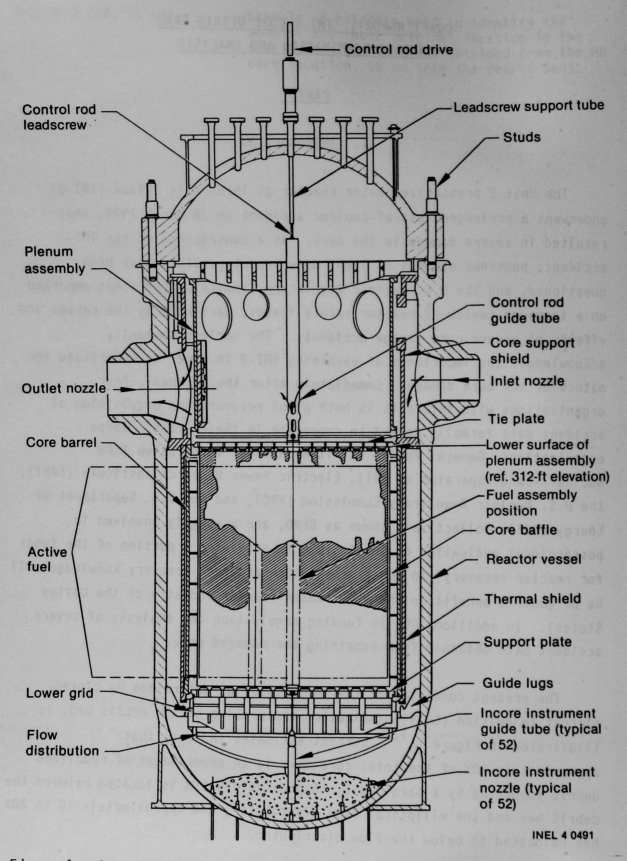


Figure 1. Schematic showing the current known condition of the TMI-2 core, as determined by CCTV and core topography examinations and debris bed probing.

The first materials obtained from the TMI-2 active fuel region were samples from the debris bed, referred to as "core debris grab samples." Six samples were obtained during September and October 1983, and five additional samples were acquired in March 1984 resulting in a total sample size of 1.37 kg out of an initial core inventory of 126 metric tons (~0.001%). The eleven samples were shipped to the Idaho National Engineering Laboratory (INEL) for examination and/or distribution to other laboratories. Ten samples were retained at INEL for examination and analysis, and one sample was shipped to the Babcock & Wilcox (B&W) Lynchburg Research Center. Of the ten samples retained at INEL, fragmented chips from seven particles were examined by Rockwell Hanford Operations (RHO) using differential thermal analysis and 22 particles were shipped to Argonne National Laboratories (East and West) for metallurgical and chemical analysis. 4,5

The primary objectives of examining the core debris grab samples include the following:

- o Supporting the plant recovery effort being performed by GPU Nuclear
- o Determining the physical, chemical, and radiochemical condition of the core
- o Providing data to support assessment of severe fuel damage codes, primarily concerning core damage progression and fission product behavior.

Results from the examination and analysis of the ten core debris grab samples retained at INEL are presented in this report. Acquisition of the samples, the examination plan, and analytical methods used in the examination are discussed in Section 2. Section 3 contains the results of the examinations, grouped into the following general categories:

- o Physical Analyses Results of the visual and photographic examinations, bulk tap density measurements, particle size distribution analysis, pyrophoricity tests, and ferromagnetic material content determination.
- o Metallurgical Analyses Analyses performed to determine visual appearance and elemental composition (on a microscale) of selected particles from the core debris materials. Estimates were made of maximum temperatures experienced during the accident. Discrete particles of debris were examined using optical metallography, scanning electron microscopy with energy dispersive x-ray spectroscopy (SEM/EDS), and scanning Auger spectroscopy (SAS).
- o Chemical Analyses Chemical analysis [e.g., inductively coupled plasma spectroscopy (ICP)] was used to measure quantitatively the elemental composition of the core debris grab samples.
- o Radiochemical Analyses Radiochemical analysis techniques, including gamma spectroscopy, neutron activation analysis, and beta emitter analysis (⁹⁰Sr), were used to measure the radionuclide concentrations of the core debris grab samples and provide information on fission product behavior. In addition, a leaching analysis (cesium release and settling test) was performed to determine radionuclide release caused by crushing the core materials.

Section 4 presents and evaluation of the condition of the TMI-2 core with regard to temperatures, core relocation, and radionuclide behavior, based on the information presented in this study and information available from other examination activities conducted to date. Conclusions and observations resulting from the examinations are presented in Section 5.

Due to its large volume, this report has been divided into two parts. Part 1 contains the main body of the report, and Part 2 contains
Appendixes A through H.

2. SAMPLE ACQUISITION, EXAMINATION PLAN, AND ANALYTICAL METHODS

This section discusses acquisition of the core debris grab samples, and describes the examination plan and analytical methods used in the examination.

2.1 Sample Acquisition

A series of CCTV inspections were made of the TMI-2 core in 1982 by inserting a miniature television camera into the upper portion of the active fuel region through leadscrew support tube and control rod guide tube openings (References 1 and 2). Three core locations (H8, E9, and B8) were chosen for initial inspection. The inspections indicated the presence of a void extending axially approximately 160 cm into the active fuel region and radially, in some areas, to the core former wall. The void space comprised approximately 25% of the original core region. Fuel assemblies were not visible in the central region of the core. Instead, there existed a bed of rubblized core material.

After the third CCTV inspection on 12 August 1982, the debris bed was probed by lowering a 1.3-cm-diameter stainless steel rod through the H8 and E9 leadscrew support tube and control rod guide tube openings. The point at which the rod touched the debris was measured. Then, the rod was rotated and allowed to penetrate the debris under its own weight (~13.5 kg). The rod penetrated easily to a depth of approximately 36 cm at both the H8 and E9 locations. The results of the probing prompted GPU Nuclear and EG&G Idaho, Inc., to obtain samples of the particulate debris from various depths. Three sampling depths were chosen: surface of the debris bed, and 8 cm and 56 cm into the bed. The H8 and E9 core locations and pre-accident locations of the control assemblies, burnable poison assemblies, and in-core instrumentation are shown in Figure 2.

Two different sampling devices were designed and fabricated to extract material from the debris bed. A clamshell type tool was used to obtain samples from the surface of the debris bed, and a rotating tube device was

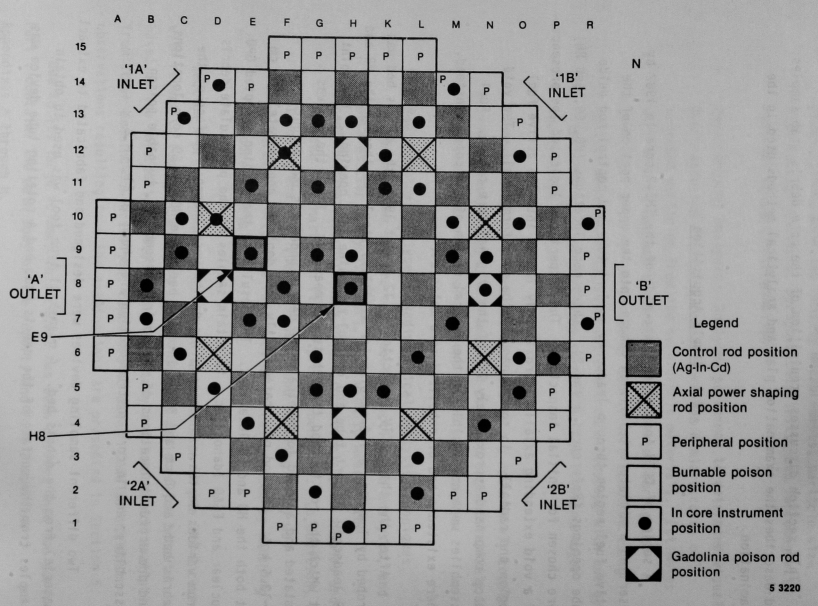


Figure 2. Schematic showing the pre-accident fuel assembly and instrumentation arrangement of the TMI-2 reactor.

used to obtain the subsurface samples. Each device was designed to hold approximately 33 cm³ of particulate debris. The sampling method and devices are illustrated in Figure 3.

The first group of six samples of particulate debris (identified as Samples 1 through 6) were obtained from the debris bed at the H8 and E9 core locations during September and October 1983. During initial probing with the sampling device, a hard stop (crust) was encountered approximately 36 cm into the debris bed at both the H8 and E9 locations. The crust was penetrated using manual force on the sampling device. [Note: A similar crust was encountered during subsequent debris bed probing at 15 other locations.] After removal, the six samples were shipped to EG&G Idaho at INEL. Subsequently, Sample 2 was shipped to the B&W Lynchburg Research Center for examination and analysis.

After acquisition of the first group of six samples, plans were made to obtain additional samples from deeper into the debris bed at the two previously sampled locations (H8 and E9). The sampling apparatus was modified to allow deeper penetration into the bed, and five samples (identified as Samples 7 through 11) were obtained in late March 1984. The maximum depth the sampling device would penetrate into the debris bed was 77 cm at the H8 location and 94 cm at E9. At the H8 (77 cm) depth, a hard stop was encountered. At the E9 (94 cm) depth, there was a friction stop encountered rather than a hard stop. The second group of five samples also was shipped to INEL for examination.

The locations of the eleven samples in the TMI-2 core^a are shown in Figure 4. The locations of the samples, both as depth into the debris bed and distance from the bottom of the plenum assembly, given in both inches and centimeters, are listed in Table 1.

a. Throughout this report, where the original location of the sample in the debris bed is pertinent to the discussion, the core location and depth of the sample in the debris bed are listed in parenthesis after the sample number [e.g., Sample 3 (H8, 56 cm)].

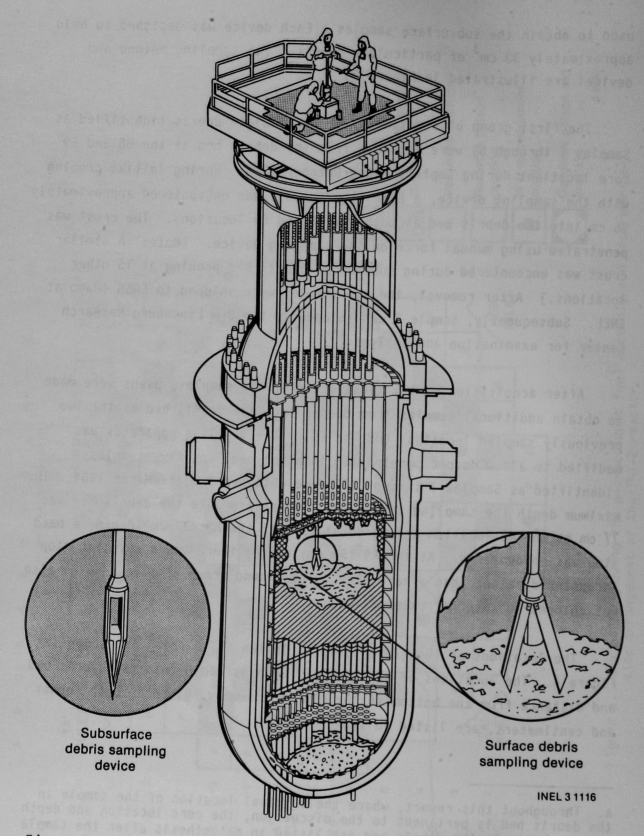


Figure 3. Schematic showing the sampling method used to obtain the TMI-2 core debris grab samples.

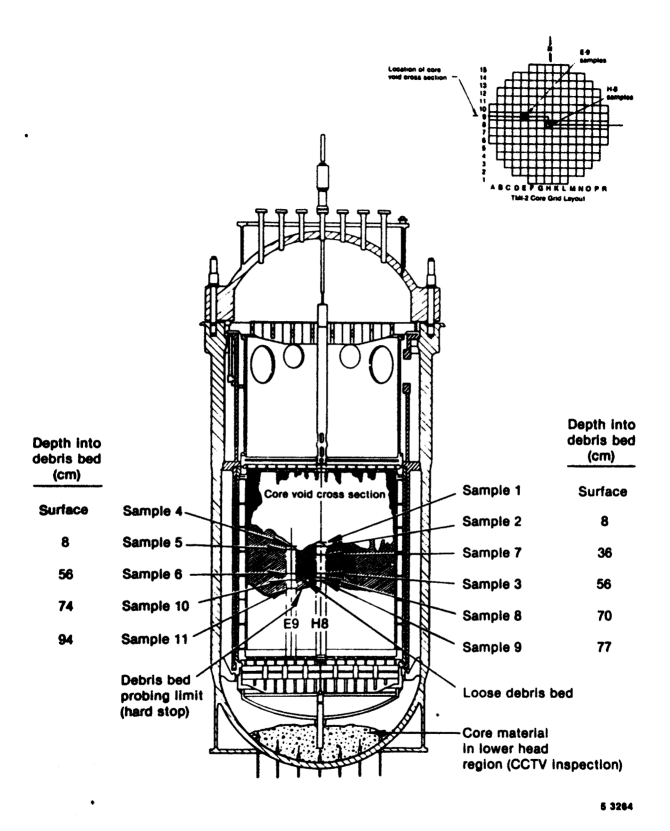


Figure 4. Locations of core debris grab samples in the TMI-2 core.

TABLE 1. SAMPLING LOCATIONS OF CORE DEBRIS GRAB SAMPLES IN THE TMI-2 CORE

Sample	Core Location	Depth into Debris Bed (in.)	Depth into Debris Bed (cm)	Distance from Bottom of Plenum Assembly ^a (in.)	Distance from Bottom of Plenum Assembly (cm)
1	Н8,	Surface	Surface	60	152
2	Н8	3	8	63	160
3	Н8	22	56	82	208
4	E9	Surface	Surface	63	160
5	E9	3	8	66	168
6	E9	22	56	85	216
7	Н8	14	36	74	188
8	Н8	27.5	70	87.5	222
9	H8	30.5	77	90.5	230
10	E9	29	74	92	234
iĭ	E9	37	94	100	254

a. The bottom of the plenum assembly is the 312-ft reference elevation in the reactor vessel design.

2.2 Examination Plan

2.2.1 Objectives

Acquisition and examination of the core debris grab samples is one of the tasks outlined in the TMI-2 Core Examination Plan which describes the technical/scientific data that should be acquired from examining the TMI-2 core. The general objectives of the examination include (a) supporting the plant recovery, (b) determining the condition of the debris bed, and (c) providing data to support assessment of severe fuel damage computer codes. Using the general objectives as a guide, an examination plan was developed which would produce data to address the following specific informational needs:

- o The physical form of the core debris (particle size, shape, morphology, origin, etc.)
- o Pyrophoric potential of materials present in the debris
- o Chemical form and composition of the various types of particles and peak temperatures to which those materials were exposed
- o Chemical form of the debris (fuel, cladding, control material, structural material, reaction product, etc.)
- o Identity and quantity of fission products retained in the debris
- o Present location of core materials
- o Leaching rates of radionuclides from existing and freshly created surfaces
- O Unanticipated defueling concerns (filtration properties, settling rates, etc.).

The information is important in planning the defueling of TMI-2. The physical form of the debris (particle size and structure) is important because small sized particles may be suspended during defueling and cause cloudiness of the water, thus obstructing operator vision. Particle size distribution data will help determine the types of filters, cyclones, etc.. needed to clean the water and remove the debris during defueling. The presence of pyrophoric materials may indicate that even larger concentrations of those materials exist below the loose debris, which must be considered during defueling and shipping of the TMI-2 core debris. Characteristics of the physical state of the debris particles (e.g., presence of prior molten materials) may provide a clue as to the nature of the core material underneath the loose debris layer. The physical and mechanical properties of the core debris materials will influence the design of tools and methods for defueling. The retained fission product content of the debris also is important because it represents a potential radiological hazard that must be controlled. The rate at which radioisotopes leach from the debris will affect the level of radioactivity in the water during defueling.

Assessment of severe fuel damage codes is important in predicting overall reactor safety. Data obtained from the damaged TMI-2 core will be the most extensive full-scale data available for code benchmarking. Information on peak temperatures, fission product transport and retention, and final physical state can be compared directly with code predictions. The TMI-2 accident scenario was inferred from a combination of code calculations, analysis of core materials, evaluation of data from on-line instrumentation, and results from other severe accident research.

2.2.2 Examination Plan

The high radiation fields (up to 100 rad/h) associated with the samples necessitated analyzing only portions of the samples in some cases. Most chemical and radiochemical analysis techniques are limited to relatively low radiation fields (<400 mrad/h). It was realized that by reducing the quantity of material analyzed to attain acceptable radiation

levels, the small portions might not be representative of the entire sample. Therefore, a sampling and analysis plan was developed which included analyzing entire samples, as well as small portions.

A flow diagram of the plan developed for examining the core debris grab samples is shown in Figure 5. The same information is presented in Table 2, except the examinations are organized according to the general category of examination (physical, metallurgical, chemical, and radiochemical). Each bulk sample was examined visually, photographed, dried, weighed, measured for bulk tap density, and steved into particle size fractions (up to ten size fractions with particle diameters ranging from less than 20 to greater than 4000 µm). Individual particles were selected from the larger (> $1000 \mu m$) particle size fractions. An aliquot (a representative portion of a homogeneous particle size fraction) was removed from each smaller (<1000 um) size fraction. The individual particles and aliquots were subjected to a variety of analyses (see Figure 5). Ferromagnetic material content examinations were performed on Sample 6, and pyrophoricity tests were performed on portions of Samples 1. 3. and 6. Approximately one-third of each particle size fraction was recombined to approximate the original bulk sample (called the "recombined bulk sample"). The recombined bulk samples were dissolved in acid solutions and analyzed to determine their average elemental and radiochemical compositions. The remainder of the particle size fractions for Sample 6 were subdivided, and a second recombined bulk sample was formed. Cesium release and settling tests were performed on this second recombined bulk sample.

2.3 Analytical Methods

This subsection summarizes the analytical methods used in examining the core debris grab samples. Brief descriptions of individual examination techniques are contained in Appendix A. The methods used for analyzing the core debris grab samples were, in most instances, standard laboratory techniques, with modifications to adapt them to the physical characteristics of the sample material and high radiation levels.

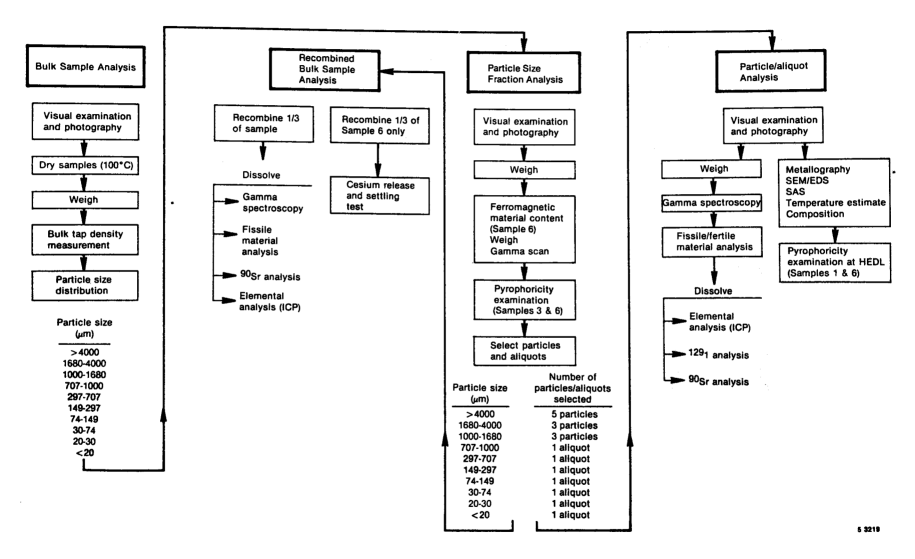


Figure 5. Flow diagram showing the examinations performed on the TMI-2 core debris grab samples (typical for each sample).

TABLE 2. EXAMINATIONS PERFORMED ON THE TMI-2 CORE DEBRIS GRAB SAMPLES

Examinations	Bulk Sample	Particles and <u>Aliquots</u>	Recombined ^a Bulk Sample
Physical	 		-
Visual and photographic	X	X	
Weight	X	X	X
Bulk tap density	X		
Particle size distribution	X		
Ferromagnetic material content	X		
Pyrophoricity	••	X	
Metallurgical			
Optical metallography		хþ	••
Scanning electron microscopy (SEM)/ energy dispersive x-ray spectroscopy (EDS)	••	Хр	
Scanning Auger spectroscopy (SAS)		Χp	~ ~
Chemical			
Inductively coupled plasma spectroscopy (ICP)		X	X
Radiochemical			
Gamma spectroscopy		X	X
Neutron activation/delayed neutron analysis for fissile fertile material content		X	χс
1291 analysis		X	***
90Sr analysis		X	X
Cesium release and settling test			X

a. After particle size analysis of each bulk sample, approximately one-third of each particle size fraction was weighed, then recombined to approximate the original sample. Hence the term "recombined bulk sample."

b. Metallurgical examinations were performed only on selected particles larger than 1000 $\mu \text{m}_{\text{-}}$

c. Fertile material content analysis was not performed on the recombined bulk samples.

2.3.1 Physical Examinations

Visual and photographic examinations were performed during unpackaging and throughout the handling operations of the bulk samples and particle size fractions to determine the sizes, shapes, and types of particles in the samples. The samples were dried at 100°C and weighed. The bulk tap density of each sample was measured by placing the sample in a graduated beaker, tapping the beaker to compact the material, measuring the volume, and dividing the sample weight (g) by the volume (cm³). The bulk tap density measurement has a relatively high (~40%) uncertainty but is fast and easy to perform. GPU Nuclear requested this examination to obtain an approximation of the debris bed density. The bulk tap density measurements results include all uncertainties resulting from voids, uneven packing, and irregular shapes of particles. A more precise (wet) density measurement was not performed because the degree of accuracy associated with that technique was not required.

The particle size distribution was determined for each bulk sample. using a column of progressively smaller sieves. New sieves were used for each sample to prevent cross contamination. After transferring (dry) the bulk sample to the sieve column, the grab sample was agitated for several minutes to segregate it into particle size fractions according to sieve aperture size. Two of the core debris grab samples (Samples 4 and 5) were not subjected to a complete particle size analysis because they are composed principally of large (>1000 µm) particles. The sieve fractions containing large (>1000 µm) particles were transferred directly to storage containers. The smaller (<1000 µm) particle size fractions were wet sieved (using a freon wash) and transferred to storage containers. Wet sieving was performed to reduce the loss of particles by aerosol transport or adherence to sieve surfaces. Each particle size fraction was examined visually, photographed, and weighed. The ferromagnetic material in each particle size fraction of Sample 6 was removed, weighed, and gamma scanned. Pyrophoricity tests were performed at INEL on material from all particle size fractions from 30 to 4000 µm for Samples 3 and 6. RHO also performed differential thermal analysis tests on fragmented chips from seven particles from Samples 1 and 6. The particle sizing, ferromagnetic determination, and pyrophoricity tests were performed at the request of GPU Nuclear to support the plant recovery effort.

After completing the physical examinations of the bulk samples, discrete particles were removed from the larger (>1000 µm) size fractions for individual examination, and aliquots were obtained from the smaller (<1000 µm) size fractions. The remainder of the sample material was temporarily stored, pending further examinations, should they be required. Selection of discrete particles was not done on a representative basis; that is, particles were chosen based on specific characteristics of interest (shininess, surface characteristics, pieces of cladding, fuel pellets, etc.). The discrete particles and aliquots were examined visually, photographed, and weighed. Portions of each particle or aliquot were removed and encapsulated in preweighed aluminum cylinders in preparation for radiochemical examination.

2.3.2 Metallurgical Examinations

The large particles generally could be visually classified in one of five categories: fuel; cladding; foamy/porous, prior molten; composite fuel/prior molten; or metallic. Particles from each category were examined metallurgically. Prior molten fuel material was identified by its foamy, glazed, or layered appearance; structural material was identified by its metallic (shiny) appearance. The particles selected for examination were not homogeneous; thus portions removed earlier from particles for differential thermal analysis at RHO and radiochemical analyses were not necessarily representative of the portion examined metallurgically.

Initially, seven particles were selected from Samples 1 and 6 for metallurgical analysis at INEL. Subsequently, 22 additional particles were selected from Samples 1 and 3 through 11 for metallurgical analysis at Argonne National Laboratories. The 29 particles were mounted in conductive Bakelite to facilitate later analysis by SEM. They were ground, to expose the surface to be examined, and polished. Photographs were taken at

magnifications from 15 to 500X. Most particles were etched using a fuel or zircaloy etchant and rephotographed. After optical metallography, 27 particles were examined by SEM/EDS. Fourteen of those particles were selected because of their unusual microstructural compositions and were further examined by SAS.

Microstructural information was obtained from the optical photomicrographs. The volume fractions and distribution of different phases and transformation structures were identified to obtain information on the thermal histories of the particles. In addition, general compositional information was obtained based on color and reflectivity of the material.

Additional microstructural information was obtained at magnifications up to 2000X, using SEM/EDS analysis. Phase and elemental distributions were obtained from the backscatter electron image, and elemental identification was obtained from EDS for elements with atomic numbers down to 12 (magnesium).

SAS was used to obtain quantitative elemental data for elements with atomic numbers down to 5 (boron). In particular, this analysis was able to quantitatively measure the oxygen content, which is critical in determining the melting temperature of U, Zr, and O mixtures.

2.3.3 Chemical and Radiochemical Examinations

Portions of selected particles from the larger (>1000 μ m) particle size fractions and aliquots from the smaller (<1000 μ m) particle size fractions from each bulk sample were analyzed by gamma spectroscopy and delayed neutron analysis to determine the radionuclide and fissile/fertile material contents. Those portions then were dissolved for subsequent chemical and radiochemical analyses. The solutions containing the dissolved core debris materials were divided so that portions could be analyzed using both radiochemical and standard chemical techniques. All decay activity data are corrected to April 1, 1984. After dissolution,

fertile material analysis could not be performed on the liquid samples because of the relatively high hydrogen content in the solutions, which would interfere with the fertile material measurements.

Initial attempts at dissolving the core debris material indicated that much of it was insoluble using normal dissolution methods (HNO $_3$ + HF). Therefore, a pyrosulfate fusion technique was modified and produced satisfactory results. The dissolution was performed in a closed system to contain the volatile 129 I and Te (Figure 6). Stable carrier I and 131 I tracer were added to the fusion to determine losses of 129 I during the dissolution process. To evaluate the loss of Te in the dissolution, several experiments were performed using 125 Te as a tracer. Experiments indicated retention of more than 80% of the 125 Te tracer in the nonvolatile portion of the dissolved material. The rest of the 125 Te tracer (~20%) was probably lost by plateout on the glassware.

The 129 I content in the volatile fraction of the dissolved material was determined using neutron activation. Subsequent gamma spectroscopy of the material determined the amount of 131 I tracer and 130 I (an activation product of 129 I).

The 90 Sr content was determined by placing the nonvolatile portion of the dissolved material in solution in water and precipitating the 90 Sr and Sr carrier out as a carbonate. The 90 Sr fraction was filtered and analyzed using the standard analysis method discussed in Appendix A.

Generally, all other elements remained in the nonvolatile portion. Inductively coupled plasma spectroscopy (ICP) was used to analyze for the following elements: Ag, Al, B, Cd, Cr, Cu, Fe, Gd, In, Mn, Mo, Ni, Nb, Si, Sn, Te, U, and Zr. These elements plus oxygen are the principal elemental

a. Volatility, as used in this section of the report, applies to dissolution and subsequent gasification of elemental components for which analyses were performed. Because of the temperature and acidity conditions during the dissolution, the volatile materials of interest would become gaseous and dissociate from the solution.

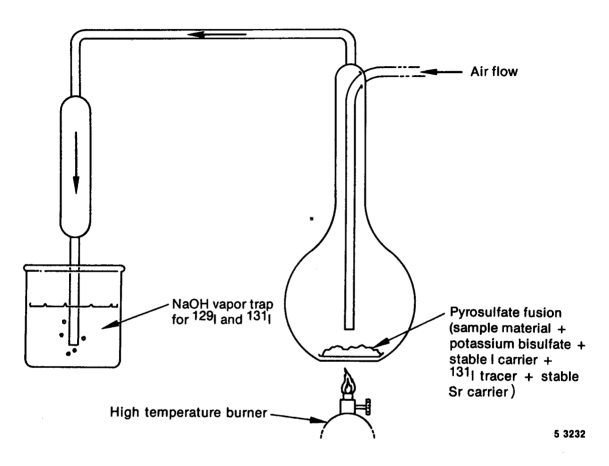


Figure 6. Schematic of the pyrosulfate fusion technique used to dissolve particles and aliquots from the core debris grab samples.

constituents present in the core. To evaluate losses of these constituents from the pyrosulfate fusion dissolution and subsequent separations, the U content measured by ICP was compared with the U content measured by delayed neutron analysis of the material before dissolution. Similar results (within 10 to 15%) were obtained, indicating that little material was lost For most samples during the dissolution and separations. The ICP results were used in determining sample compositions and ratios of the elements.

Each recombined bulk sample was dissolved and analyzed to measure its average elemental and radiochemical concentration. Dissolution of the recombined bulk samples was difficult because of the relatively large (up to 50 g) mass of highly insoluble debris. The dissolution was performed in a hot cell because of the high radiation fields (up to 100 rad/h, β/γ). Safety concerns prevented using the pyrosulfate fusion technique in a hot cell because of the potential for fire or explosion. Therefore, a sequential dissolution was performed using acids of lesser strengths. The dissolutions were performed in a closed quartz system similar to Figure 6, but at lower temperatures using a hot plate instead of a high temperature burner. The acids and sequence of use were: (a) $6\underline{M}$ HNO $_3$, (b) $3\underline{M}$ HNO $_3$ + $1\underline{M}$ HF, and (c) aqua regia. Dissolution of each recombined bulk sample required 3 to 4 days, with several additions of HC1 to the aqua regia solution.

The resulting volume of solutions for each recombined bulk sample ranged from 300 mL to 1 L, with specific activities up to 200 mrad/h per mL of solution. Small (~10 mL) portions of each solution were removed, diluted, and analyzed using the techniques described previously for the particles and aliquots. No 129 I analysis was performed because the sequential dissolution made use of stable carrier I and 131 tracer unfeasible.

3 FXAMINATION RESULTS/DISCUSSION

This section presents results of the physical, metallurgical, chemical, and radiochemical examinations of the TMI-2 core debris grab samples, as described in Section 2. Principal results include the following:

- Most of the particles examined contain regions of prior molten U-Zr-0, indicating that peak temperatures greater than 2800 K were reached in the core. Also, there are a few prior molten particles that are almost pure $U0_2$, suggesting temperatures up to 3100 K.
- The regions of prior molten U-Zr-O usually contain at least traces of some structural materials (Al, Cr, Fe, and Ni). These elements often are found at grain boundaries or in voids. For some particles, structural materials are a significant fraction of the particle. Control rod materials (Ag, In, and Cd) are not as prevalent as structural materials.
- o Ag, a control rod material, apparently had less of a tendency to mix or interact with fuel rod materials (Zr and U) than did structural materials. Significant depletion of Ag (~90%) from the samples was observed.
- o The data suggest the occurrence of prolonged candling sequences or multiple temperature excursions during the accident.
- Evidence was found of the flow of molten metallic U and Zr inside ballooned cladding.
- Significant Zr depletion (\leq 50%) has occurred from the majority of the samples analyzed, indicating significant relocation of Zr from the debris bed.

- Evidence of significant relocation/mixing of core materials is inferred from the distribution of Gd and the U enrichments.
- o Al and Te appear to have concentrated at the surface of the debris bed.
- o The radionuclides are retained in the samples in the approximate order of their volatilities.
- o Some evidence was found which indicates a correlation between Ni content and 125 Sb and 106 Ru concentrations.

The results are discussed in detail in the following subsections.

3.1 Physical Analyses

This section describes the results of the visual and photographic examinations, weighing, bulk tap density measurements, particle size and weight frequency distribution analyses, categorization of particle types, and ferromagnetic and pyrophoricity tests performed on the 10 samples. These examinations provided the first information on the condition and composition of the core debris material. It was determined through these tests that the particles range from approximately 10 millimeters to a few micrometers in size. The particles were categorized visually by type. There are five general types of particles ranging from basically unaffected segments of fuel pellets and cladding to prior molten material. The smaller (<1000 µm) sized particles contain about 10% ferromagnetic material.

The results of the physical examinations were used to help develop the follow-on examination plan and select the types and quantities of materials for the subsequent metallurgical, chemical, and radiochemical examinations.

3.1.1 Visual and Photographic Examinations and Weights of Bulk Samples

As the sampling tools were opened and the contents removed, visual examination was conducted and photographs were obtained of each sample. Photographs of each bulk sample are presented in Appendix B. Overall description of the ten samples, including their original locations in the reactor, weights, gross gamma radiation levels, and visual characteristics as first examined are presented in Table 3.

As the sampling tools were opened, most of the samples were observed to be stratified according to particle size (the larger particles were at the top and the smaller particles at the bottom of the sampling tool chamber). Because most of the subsurface samples had this type of stratification, it was concluded that the smaller particles settled to the bottom of the sampling tools as a result of the sampling technique and/or handling and shipping operations. The debris bed most probably would not have had a similar stratified distribution at each of the several depths sampled.

3.1.2 Bulk Tap Density

Dry bulk tap density measurements were made for Samples 1, 3, 6, 9, 10, and 11 (Table 4). The average of the densities from the first group of samples (Samples 1, 3, and 6) is 30% less than that from the second group (Samples 9, 10, and 11). The particle size distribution analysis performed subsequently (see Section 3.1.3) revealed that Samples 1, 3, and 6 have the largest particle size fraction of material in the 1680 to 4000- μ m size range, whereas Samples 9, 10, and 11 show a bimodal distribution, with peaks at the 1680 to 4000- μ m and 297 to 707- μ m sizes. The greater weight percent of small particles in Samples 9, 10, and 11 probably allowed more efficient packing and, therefore, caused the higher bulk densities.

TABLE'3. RESULTS OF VISUAL AND PHOTOGRAPHIC EXAMINATIONS, WEIGHING, AND GROSS RADIATION SURVEY OF THE TEN BULK SAMPLES

Sample	Sampling Tool	TMI-2 Core	Depth of Sample in Debris Bed (cm)	Sample Weight	Gamma Radiation Level ^a (rad)	Visual Characteristics ^b
1	Clamshell	нв	Surface	70.88	16 ^C	A pile of very black, damp debris with a fairly wide range of particle sizes (up to 10 mm); several rounded surfaces; sporadic rust color throughout.
3	Rotating tube	Н8	56	152.71	36c	Very black, slightly damp debris with wide range of particle sizes (up to 5 mm); small chunks to fine debris; similar to Sample 1.
4	Clamshell	£ 9	Surface	16.59	3c	Thirteen major chunks of black, dry debris with rust colored sides; basically sharp edges with one or two chunks having rounded edges; dimensions range from 5 to 10 mm.
5	Rotating tube	E9	8	90.96	18 ^C	Similar to Sample 4 with the following differences: many more pieces, greater size range (1 to 10 mm), some surfaces more reflective; very dry.
6	Rotating tube	E9	56	140.73	36c	Very black, slightly damp debris; small chunks to fine debris; some pieces blackish gray; a few pieces resemble metal shards similar to Sample 3.
7	Rotating tube	н8	36	135.86	20 ^d	A wide range of particle sizes (up to 10 mm); porous surfaces; sporadic rust areas.
8	Rotating tube	Н8	70	152.76	22 ^d	A wide range of particle sizes (up to 5 mm); smooth, angular surfaces; pieces of cladding.
9	Rotating tube	H8	77	153.08	25 d	Very fine, dark debris with only a few larger rounded chunks and metallic particles.
10	Rotating tube	E9	74	173.90	30 ^d	A wide range of particle sizes (up to 10 mm); rounded, angular surfaces.
11	Rotating tube	E9	94	148.75	20 ^d	Similar to Sample 10 with the following distinctions: a larger fraction of particles resemble fuel pieces, more smaller chunks.

a. Radiation levels measured using a teletector probe on the exposed sample.

b. All samples appeared to be of a loose granular nature with some sharp or rounded edges.

c. Reading at 2.5 cm.

d. Reading at 5 cm.

TABLE 4. BULK TAP DENSITIES OF THE TMI-2 CORE DEBRIS GRAB SAMPLES

<u>Sample</u>	TMI-2 Core Location	Depth of Sample in Debris Bed (cm)	Bulk Tap Density ^a (g/cm ³)
1 3 6	H8 H8 E9	Surface 56 56	3.5 3.8 3.5
			3.6 ^b
9 10 11	H8 E9 E9	77 74 94	5.0 5.5 5.0
			5.2°

a. Bulk tap densities were not measured for Samples 4, 5, 7, and 8.

b. Average of Samples 1, 3, and 6.

c. Average of Samples 9, 10, and 11.

3.1.3 Particle Size Distribution

The particle size distribution of the samples was determined using the method described in Section 2.3.1. The particle size fractions were weighed and photographed (see Appendix B). Results of the particle size distribution analyses are presented in Table 5. Also, the cumulative weight distributions of each sample and the combined samples are shown in Figures 7 and 8. The particle size distribution as a function of depth into the debris bed is shown in Figure 9.

Five samples from the H8 core location (Samples 1, 7, 3, 8, and 9, in order of depth into the debris bed) were examined at INEL. The particle size distributions of Samples 1, 7, and 8 are similar in that 81 to 86 wt% of the sample material is larger than 1000 µm in size, and the 1680 to 4000-um particle size fraction contains the most material. Sample 3 had 90 wt% of its material larger than 1000 um in size; however, the largest amount of material was in the greater than 4000-um fraction. was obtained about 20 cm below where a breakable crust had been contacted during the sampling process, which may account for its greater weight percent of larger sized particles. Sample 9 was the deepest sample obtained from the H8 location. During sampling, a hard stop was encountered by the sampling tool at 77 cm into the debris bed. Only 59 wt% of the material was larger than 1000 µm in size, indicating either some setlling of the fine material occurred in the debris bed or washout of the fine material occurred from the upper layers. Sample 9 also showed a bimodal distribution of particles, with the major peak at the 1680 to 4000-um size fraction and a minor peak at the 300 to 710-um size fraction. The bimodal distribution indicates a more efficient packing and subsequent higher density for this region of the debris bed.

Five samples (Samples 4, 5, 6, 10, and 11, in order of depth into the debris bed) were removed from the E9 location (about mid-radius in the core). The first two samples (Samples 4 and 5) contain only large (>1000 µm) particles, which correlated with observations made during the CCTV inspection at this location. Sample 6 (E9, 56 cm) was similar to

TABLE 5. RESULTS OF THE PARTICLE SIZE ANALYSIS OF THE TMI-2 CORE DEBRIS GRAB SAMPLES

Particle Size Fraction	Samp	le l	Samp1	e 3	Samp	le 4ª	Samp	ole 5 ^b	Samp	le 6	Samp	le 6 ^C
(μm)	<u>(e)</u>	(wt%)	<u>(p)</u>	(wt%)	(g)	<u>(g)</u>	(wt%)	<u>(p)</u>	(wt%)		2)
>4000	12.62	18.4	63.75	42.9	_		69.57	77.1	57.99	42.0		a
1680 to 4000	27.82	40.6	51.45	34.7			13.96	15.5	49.39	35.8	0.3	39
1000 to 1680	15.64	22.8	19.19	12.9	-		6.25	6.9	13.88	10.1	0.3	
>1000		81.8		90.5	-			99.5		87.9		:-
<1000					-		0.44	0.49				-~
707 to 1000	7.80	11.4	5.49	3.7	-				8.93	6.5	0.2	
297 to 707	3.20	4.7	6.34	4.3					5.99	4.3	0.	
149 to 297	0.87	1.3	1.27	0.86					0.97	0.70	0.0	
74 to 149	0.44	0.64	0.77	0.52	•				0.67	0.48	0.0	024
30 to 74	0.17	0.25	0.18	0.12					0.22 d	0.16	•	d d
<30	d		0.013	0.01	•				u	d	-	
Summed weight	68.56		148.45				90.22		138.04		1.	178
Initial weight	70.88		152.71		16	.59	90.96		140.73		-	
Sample loss ^e	2.32	3.3	4.26	2.8			0.74	0.8	2.69	1.9	•	
	Samp	le 7 ^f	Sampl	e 7 ^g	Samp	le 8	Samp	le 9	Sampl	e 10	Sampl	e 11
	(e)	(wt%)	<u>_(e)_</u>	(wt%)	<u>(e)</u>	(wt%)	<u>(e)</u>	(wt%)	<u>(e)</u>	(wt%)	<u>(e)</u>	(wt%)
>4000	7.38	5.46	5.48	4.04	28.39	18.61	5.69	3.73	15.93	9.32	30.58	20.59
1680 to 4000	59.44	43.98	54.14	39.89	74.55	48.86	53.81	35.26	88.03	51.49	57.44	38.67
1000 to 1680	43.29	32.03	43.70	32.20	28.91	18.95	30.63	20.07	33.56	19.63	21.20	14.27
>1000	110.11	81.47	103.32	76.13	131.85	86.42	90.40	59.23	137.52	80.44	109.22	73.53
707 to 1000	13.48	9.97	15.70	11.57	9.13	5.98	16.18	10.60	11.59	6.78	10.89	7.33
297 to 707	9.77	7.23	12.11	8.92	7.49	4.90	25.31	16.58	14.76	8.63	16.90	11.38
149 to 297	0.89	0.66	1.61	1.19	1.35	0.88	9.03	5.92	3.96	2.32	6.57	4.42
74 to 149	0.54	0.40	0.93	0.69	0.95	0.62	5.34	3.50	1.78	1.04	2.91	1.96
30 to 74	0.35	0.46	1.77	1.31	1.20	0.79	5.66	3.71	1.36	0.80	2.03	
<30 to 74	0.02	0.01	0.27	0.20	0.60	0.39	0.96	0.63				
Summed weight	135.16		135.71		152.57		152.61		170.97		148.52	
Initial weight	135.86		135.71		152.76		153.08		173.90		148.75	
Sample losse	0.70	0.52			0.19	0.12	0.47	0.31	2.93	1.68	0.23	0.16

a. Sieving was not done. Sample consisted of only large particles.

b. Sieving was limited to four sizes. Sample consisted of mostly large particles.

c. Ferromagnetic material weights. These values are subsets of their respective particle size fractions for Sample 6.

d. None detected (not measurable).

e. The sample loss defines the quantity of material lost during seiving; however, the particle size distribution of the lost material is not known.

f. After 5-min shake.

g. After 1-h shake.

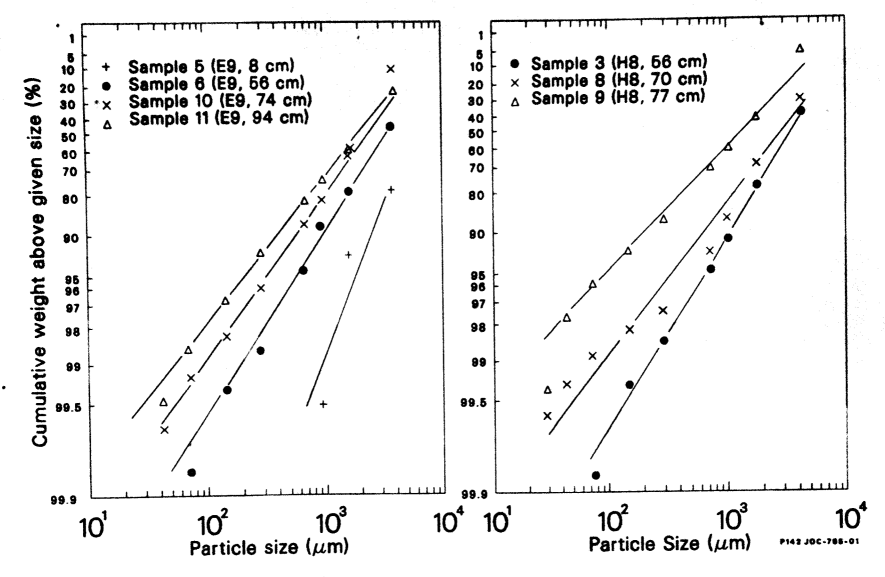


Figure 7. Plots showing the cumulative weight distribution of various core debris grab samples.

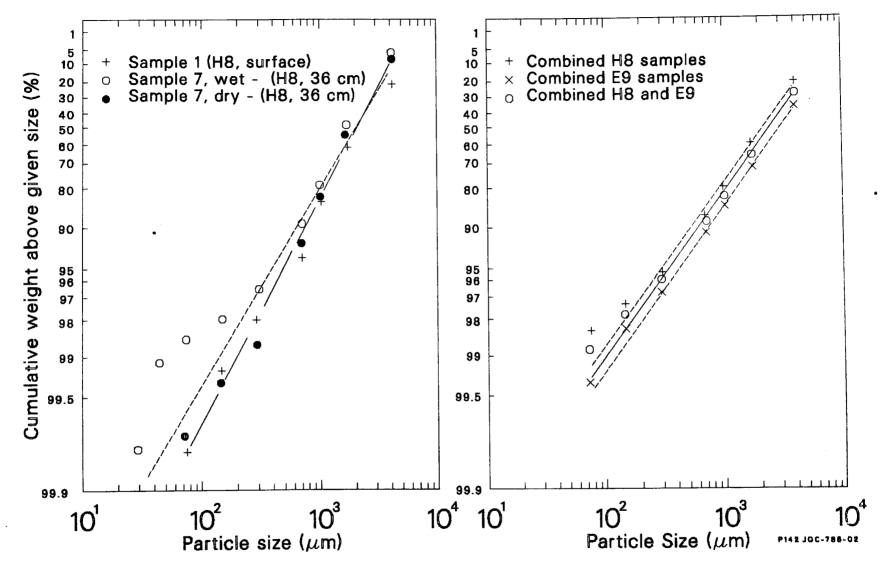


Figure 8. Plots showing the cumulative weight distribution of various core debris grab samples and of the combined samples.

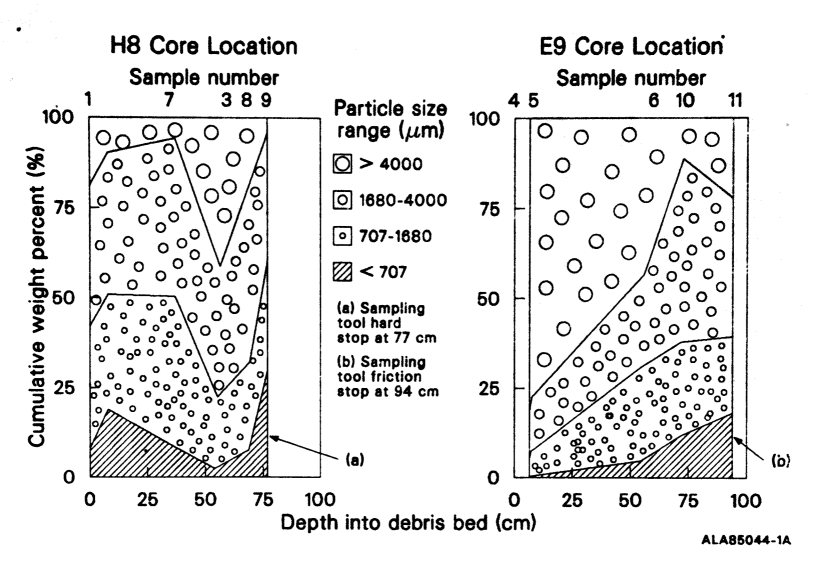


Figure 9. Plot showing the cumulative weight distribution of each core debris grab sample as a function of depth into the debris bed at the H8 and E9 core locations.

Sample 3 (H8, 56 cm) in that about 88 wt% of the material was larger than 1000 μ m, of which the largest amount was greater than 4000 μ m in size. Samples 3 and 6 were obtained from the same depth (56 cm) into the debris bed, indicating that the effect of the breakable crust may be fairly uniform across the reactor core. Samples 10 and 11 were similar to Sample 9 from the H8 position in that they exhibited a bimodal particle size distribution.

3.1.4 Weight Frequency Distribution

The weight frequency is defined as the distribution of the sample weight as a function of particle size fraction. The following weight frequency distribution information was extracted from GPU Nuclear evaluations of the particle size distribution analyses performed at INEL and B&W. The distribution of weight frequency, for each sample is shown in Figures 10 and 11.

The weight frequency, f, of particle size Dp is calculated as follows:

$$f = \frac{\Delta R}{\Delta D p} \tag{1}$$

where

f = weight frequency (%/ μ m)

ΔR = weight percent of particles belonging to the given particle size range (%)

 ΔDp = span of the given particle size range (μm).

The weight frequency was calculated with the assumption that all particles were in the 10 to $6000-\mu m$ range.

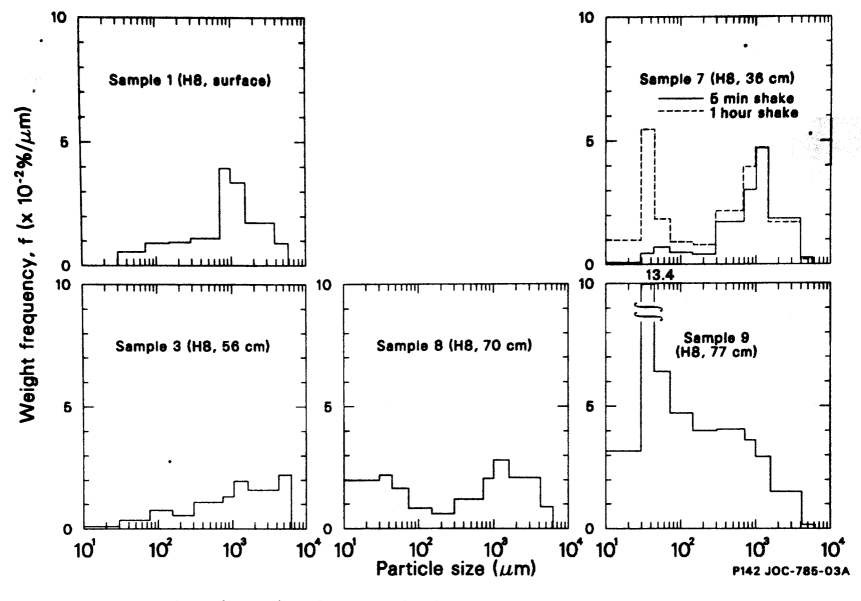


Figure 10. Weight frequency distributions of samples from the H8 core location.

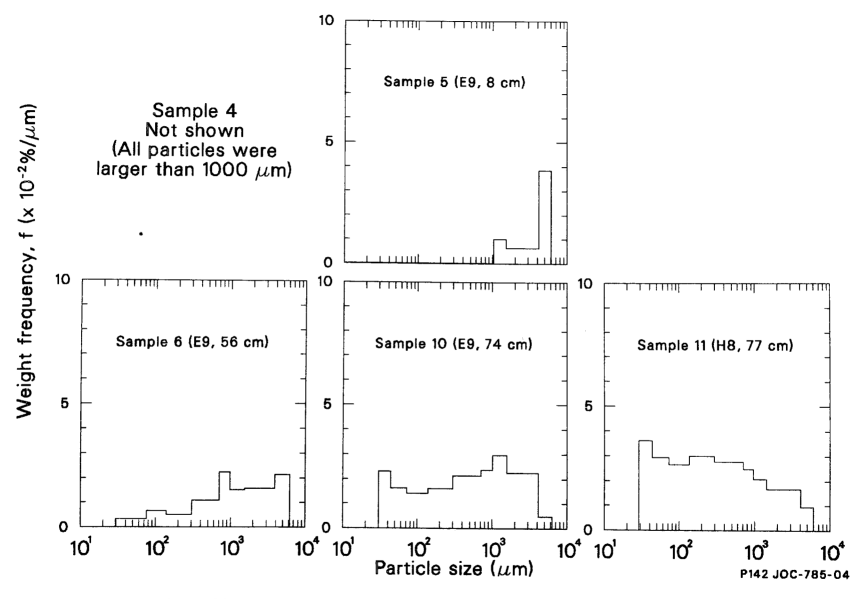


Figure 11. Weight frequency distributions of samples from the E9 core location.

The effect of agitation time during sleving for Sample 7 is shown in Figure 10. The 1-h shake caused conversion of portions of the material in the greater than 4000 μ m to 1680- μ m size ranges to the smaller size ranges, notably the 30 to 74 μ m range.

From Figure 10, the following observations were made of samples from the H8 location:

- 1. The mode sizes (particle size with the highest f value) of Samples 1 and 7 are in the range of 707 to 1680 μm .
- 2. The mode sizes of Samples 3, 8, and 9 decrease as the depth into the debris bed increases.
- 3. The weight frequency of the largest particles (i.e., greater than 4000 µm, shown as 4000 to 6000 µm in Figure 10) decreases as the depth increases for the group of Samples 1 and 7 and the group of Samples 3, 8, and 9, but it is discontinuous between Sample 7 (H8, 36 cm) and Sample 3 (H8, 56 cm).
- 4. for Sample 9 (H8, 77 cm), the 30 to 74 µm particle size range shows a very high weight frequency. This suggests that the small particles were prevented from falling deeper at the H8 location by a blockage which was felt as the hard stop during sample acquisition.

Observations 1, 2, and 3 suggest that the debris particles at the H8 location are stratified into two layers, with the boundary located between 36 cm and 56 cm below the surface of the debris bed. This is supported by the existence of a penetrable "crust-like" layer felt during sample acquisition.

From Figure 11, the following observations were made of samples from the E9 location:

- 1. The mode sizes of Samples 5, 6, 10, and 11 decreases as the depth into the debris bed increases.
- 2. No particular discontinuity is apparent between the weight frequency distributions of samples from the E9 location.

3.1.5 Particle Types

The larger particle size fractions for each sample were examined visually after particle sizing, and five general (types) categories of particles were identified based on visual appearance, including (a) apparent fuel pieces; (b) cladding chunks; (c) foamy/porous, prior molten material; (d) particles that are a composite of fuel and prior molten material; and (e) metallic appearing, prior molten particles. Based on subsequent metallography and chemical analysis of several particles from each type, the composite particles generally are assumed to be fuel pellets with a prior molten U-Zr-O material coating the surface or gluing the fuel particles together. The composite particles have rounded edges and corners and sometimes a glazed surface appearance. It is assumed that the metallic appearing, prior molten particles are probably the structural materials with lower melting temperatures. Examples of particles from each of the five types are shown in Figures 12 through 16.

In general, all particle types were found in all samples. However, some trends could be noted. Samples 1, 7, and 8 from the H8 location appeared similar visually. No easily recognizable cladding pieces were found in the largest particle size fractions of those samples, although subsequent examination did reveal some cladding pieces with fuel attached in all three samples. About 15% of the larger sized particles were recognized as fuel pieces, and the remaining material consisted of foamy, highly porous particles and composite particles. Sample 7 had a higher fraction of foamy particles than did Samples 1 and 8. Sample 3 was similar to Samples 1, 7, and 8 in that the fuel pieces composed about 15% of the large particles, however, the sample contained mostly composite particles

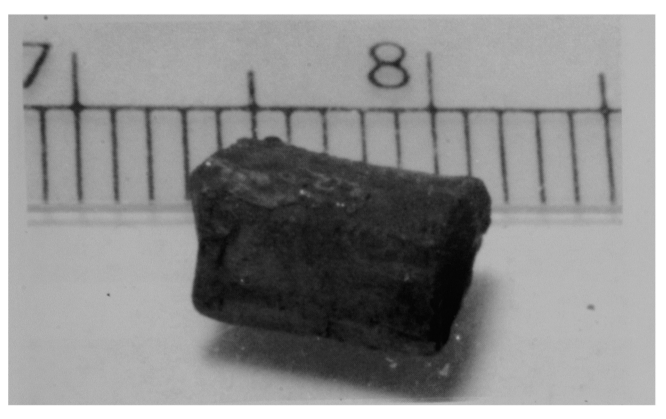
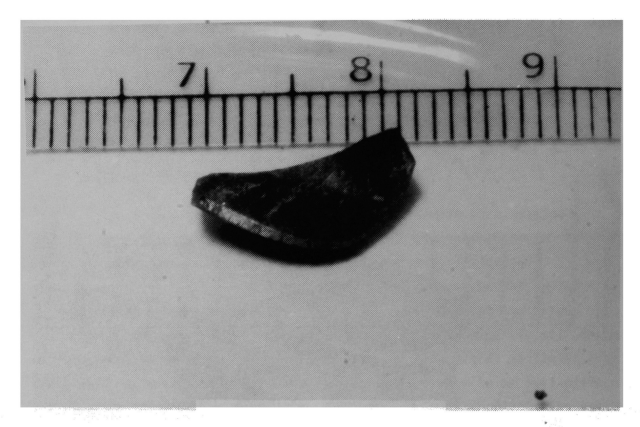


Figure 12. Photograph of Particle 1B (H8, surface) showing an example of a fuel piece.



Front view of particle

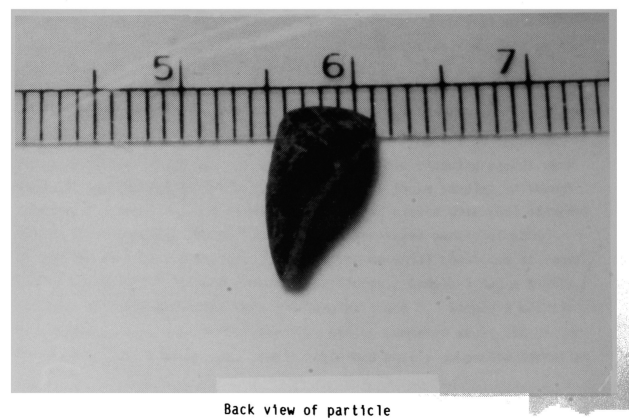


Figure 13. Photograph of Particle 6E (E9, 56 cm) showing an example of a cladding piece.

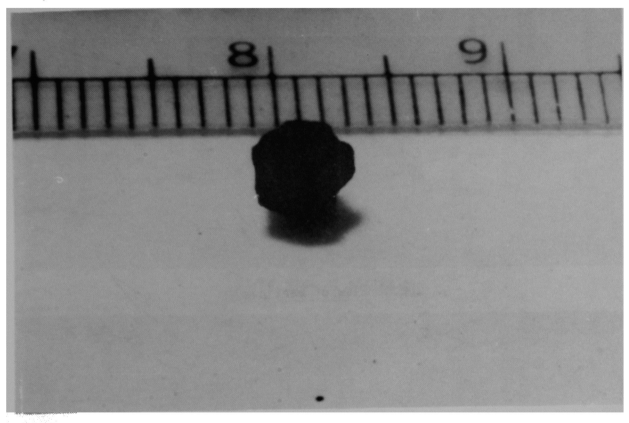
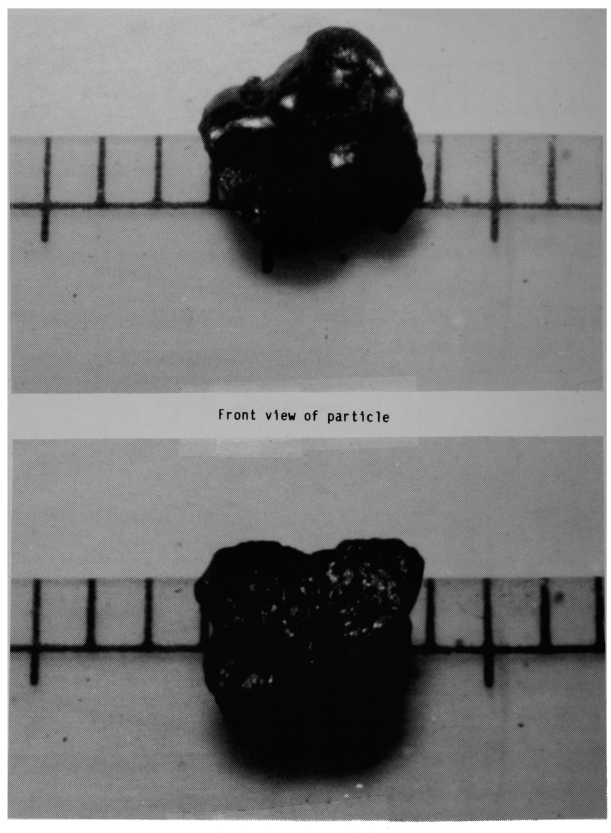
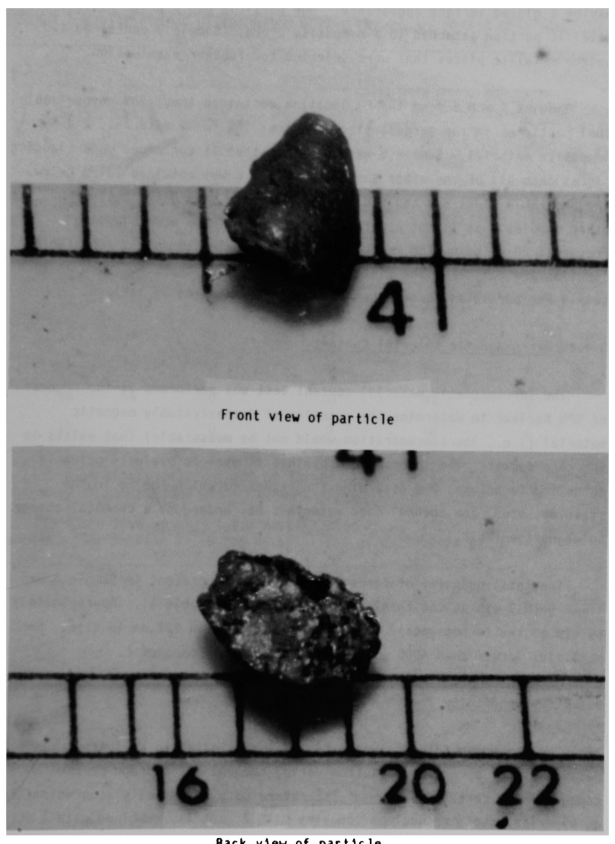


Figure 14. Photograph of Particle 1H (H8, surface) showing an example of a foamy piece.



Back view of particle

Figure 15. Photograph of Particle 8C (H8, 70 cm) showing an example of a composite piece.



Back view of particle

Figure 16. Photograph of Particle 9G (48, 77 cm) showing an example of a metallic piece.

having a glazed surface appearance. One particle had a prior molten, metallic portion attached to a composite piece. Sample 9 contained two unique metallic pieces that were selected for further examination.

Samples 4 and 5 from the E9 location contained about 30% recognizable fuel particles in the largest size fraction, 15% foamy material, and 55% composite material. Sample 6 was unique in that it contained more cladding pieces than all of the other samples. Sample 6 was obtained 20 cm below the breakable crust and had a different particle size distribution than the other samples. It is not apparent why there would be more cladding fragments 20 cm below the breakable crust. Samples 10 and 11 from the E9 location are similar. The distribution of particle types observed in each sample for particles larger than 1000µm are summarized in Table 6.

3.1.6 Ferromagnetic Material Content

The ferromagnetic material content test was performed at the request of GPU Nuclear to determine the quantity of demonstratably magnetic material (i.e., low concentration would not be measurable) that exists in the core debris. The information obtained is used to evaluate potential defueling techniques and determine the extent to which the Fe in the stainless steel and Inconel core materials has undergone a chemical change to magnetite (Fe_3O_4).

The total quantity of ferromagnetic material present in Sample 6 was 1.178 g (0.9 wt% of the total sample weight) (see Table 5). Approximately 95 wt% of the ferromagnetic material was larger than 297 μm in size. No particles larger than 4000 μm were found to be ferromagnetic.

3.1.7 Pyrophoricity Study

The phenomenon of pyrophoricity was investigated to evaluate the possibility of pyrophoric reaction during removal and storage of the damaged TMI-2 core. A review of literature on pyrophoricity (concentrating on experiences of the nuclear industry with Zr metal), combined with

TABLE 6. PARTICLE TYPE DISTRIBUTION OF THE TMI-2 CORE DEBRIS GRAB SAMPLES

			Particle Type	s ^b		
Sample	<u>Fuel</u>	Cladding	Foamy/Porous	Composite	Metallic	
1	15	0	20	65	0	
3	15	0	10	75	0	
4	30	0	15	55	0	
5	30	0	15	55	0	
6	50	20	10	10	10	
7	15	0	64	21	0	
8	12	2	27	59	0	
9	18	0	0	73	9	
10	54	0	0	46	0	
11	48	0	7	45	0	

a. Given as approximate percent of larger (>1000 $\mu m)$ sized particles. Less than 1% was rounded off to zero.

b. There are five general types of particles found in the core debris grab samples. See Section 3.1.5 for descriptions of each type.

evaluation of core material reactions at high temperature that occurred during the TMI-2 accident, indicate that pyrophoric reactions during defueling would be unlikely.

A series of ignition tests were performed at INEL on samples of core materials removed from all size fractions between 30 and 4000 µm of Samples 3 and 6. The samples were tested under both wet and dry conditions, and ignition of the material was attempted using both a Tesla Coil and a propane torch. In no case did ignition of the materials occur. 7

Differential thermal analyses were conducted by Rockwell Hanford Operations on fragmented chips from seven particles (Appendix H). The particles generally showed little thermal activity, although one particle produced a large, broad release of energy (exotherm) of 76l cal/g, spanning nearly 500 degrees, starting at about 550°C. This exotherm occurred at much higher temperatures than those observed for samples of Zr powder and partially oxidized Zr powder. If the exotherm is due to the oxidation of Zr, the Zr in the sample must have been coated with a thick non-combustible, possibly oxide, layer which protected it at lower temperatures. The results of this test are contained in Appendix H.

3.2 <u>Metallurgical Analyses</u>

Twenty-nine particles from the ten core debris grab samples retained at INEL were selected for metallurgical analyses. Twenty-two particles were studied jointly by EG&G Idaho and Argonne National Laboratories (East and West), while the remaining seven were characterized by EG&G Idaho and Westinghouse Idaho Nuclear Co. This section presents results of the optical metallography, scanning electron microscopy with energy dispersive x-ray spectroscopy (SEM/EDS), and scanning Auger spectroscopy (SAS). From these analyses, the microstructural appearances and elemental compositions of the particles were determined, estimates were made of maximum temperatures experienced during the accident, and interactions between fuel rod and non-fuel rod materials were investigated.

3.2.1 Summary of Examinations

The metallurgical analyses provided data in the form of optical pholographs, SEM backscatter and secondary electron images, x-ray spectra from the SEM/EDS examinations, and quantitative elemental compositions from the SAS analyses. The area fractions of the major components, grain sizes, and peak temperature estimates are summarized in Table 7. Element identification is based on the SEM/EDS data, except for oxygen content which is based on the SAS data, (note that not all particles were analyzed by SAS). The oxygen contents listed in Table 7 represent the ranges found in the base material (not inclusions or grain boundary phases). Area fractions of the various components were determined from visual examination of optical photographs and SEM images. Fuel grain size was determined using the line intercept method with a 2-D to 3-D conversion factor of 1.51. The grain size listed for Particle 4A was from the minimum grain size region of that particle. The peak temperature estimates shown in Table 7 are discussed in Section 3.2.2. The "other phases" referred to in Table 7 are the various materials found in grain boundaries and voids. along with the trace amounts of Fe that were sometimes found in the base material, and are discussed in Section 3.2.3. The composition of inclusions represents the average or typical inclusion (see Section 3.2.3).

Details of the metallurgical data are presented in Appendix C, including representative optical photographs and SEM images, and detailed qualitative SEM/EDS and quantitative SAS data. Appendix C also contains brief descriptions of each particle, and summarizes elemental compositions of each particle.

3.2.2 Temperature Estimates

Estimation of the temperature history is important for many aspects of the analysis of the TMI-2 accident and is required for defining the

a. M. I. Mendleson, "Average Grain Size in Polycrystalline Ceramics," <u>Journal of the American Ceramic Society, Vol. 52</u>, No. 8, August 1969, p. 443.

1ABLE 7. SUMMARY OF METALLURGICAL ANALYSIS OF THE TMI-2 CORE DEBRIS GRAB SAMPLES

į	<u>Particle</u>	Area ^a fraction of UOx	Grain Size (µm)	Area ^a Fraction <u>of {U,Zr}0x</u>	Area ^a Fraction <u>of ZrOx</u>	Area ^a Fraction of Other <u>Phases</u>	Composition of Inclusions	Area Fraction ^a of Unoxidized Metallic Regions (<u>not inclusions)</u>	Traces of Iron Found in Base Material	Oxygen Content Range of Base Material from Auger Analysis (at%)	Estimated Peak Iemperature (K)
	14	0.5	10	0.3 (U,Zr) metallic	Ίr			0.2 Zr		·	2170-2245
	18	1	12	Īr	Tr			e e e e e e e e e e e e e e e e e e e			2170-2600
	1E	0.1		0.1	0.2			0.6 Zr	Yes	~67b	2810-2 9 60
	1H			0.8		0.2α'	-			; 	>2810
	3 L			4		• -					>2850
	3M	0.4	11	0.6					Yes		>2850
46	44	0.3	>18	0.5		0.2α'	N1-Sn, Ru-T1		Yes		~2810
6	48	0.3	>18	0.7		Tra'	Ni-Sn, Ru-Tc			64-70	~2010
	4D	<u>.</u>		1			Cr-Fe-Ni-Sn				>2850
	5E	0.1	~12c	0.9		Tra	Ru-Ni-Tc-Rh		Yes	68-74	~2810
	68					. -		1.0 Zr		- •	<2245
	6C	•		0,.5	0.3	0.2ß	N1-Sn-Zr-A1- Zr <i>-</i> U	 –		63-69	2170-2960
	6D	0.9	10	0.1						~70	2170-2810
	6 £				0.2			0.8 Zr		w	~2000
	6F			0.8		Τεα'		0.2 N1 (Fe)	Yes		2170-2810
	7A			0.6		0.40'	N1-fe		Yes	-	~2810
	78	9.3	~10¢	0.2 (U,Zr) metallic	0.2			0.3 Zr			2170-2245
	7E	0.3	28	0.6		Tra',1r6	N1-Sn-Fe	0.1 N1 (Fe)	- 	65-70	~2610
	88			1		Trai	Mo-Sn			66-70	~2810
	8 C	•	***	0.6	. ·	0.2a', 0.28	Λg	• • • • • • • • • • • • • • • • • • •	Yes	62 -66	>2170

TABLE 7. (continued)

	article	Area ^a Fraction of UOX	Grain Size (um)	Area ^a Fraction of (U.Zr)Ox	Area ⁸ Fraction of ZrOx	Area ^a Fraction of Other Phases	Composition of Inclusions	Area Fraction ^a of Unoxidized Metallic Regions (not inclusions)	Traces of Iron found in Base Material	Oxygen Content Range of Base Material from Auger Analysis (at%)	Estimated Peak Temperature (K)
	8€	• •	***	0.7		0.1a 0.26	N1-Sn		Yes	70-72	-2010
	811	•	**	0.8	-	Tra	N1-Sn	0.2 Ag	Yes		>2850
	90	-	, .: +=	0.1		0.2¢	N1-Sn	0.7 Ag	Yes		~2810
	96	•	•••			0.2c	N1-Sn	0.8 Ag			>1233
	10A	0.2	28	0.8	. ••	, 	N1-Sn-Tc-Ru			65-69	~3100
	10E	0.9	10	0.1	Tr	Tra	N1-Sn	r A A g *	• • • • • • • • • • • • • • • • • • •	63-71	>2810
	10f	•		0.9	· ·	0.1a	N1-Ru-Fe	yo al	Yes	68-71	>2850
47	118	••		0.5	**	0.14 0.1c	N1-Fe-Sn-Ag- Mo	0.3 N1 (fe)			~2810
8	110	3	10	lr .	Tr	**	en e				2170-3120

Area fractions were estimated by visual examination.

Other phase compositions:

a .	_	Cr-Fe-N1	with	varying	amounts	of	U-Zr.

Al-Cr-Fe-Nt with varying amounts of U-Zr. High Al with Zr-U some Fe-Cr.

Austra Contest

Oxygen content refers to attached (U,Zr)O2 only.

c. Problems were encountered in measuring these grain sizes.

Tr. Contains trace amounts of these materials.

Different (from base) U-Zr compositions.

Separate Cr-fe and N1-fe bearing phases.

N1-Sn with some Ag-Fe (metallic phase).

accident scenario. For example, estimates of fission product release are extremely temperature dependent, with highly volatile fission products released at lower fuel temperatures (<2100 K) and less volatile fission products released as temperatures approach UO₂ melting. While it is not possible to determine temperature history from examining the core debris grab samples, peak temperatures experienced by the material can be estimated in many cases. These peak temperatures and other TMI system response data can be used in conjunction with accident analysis codes to establish a temperature history.

Peak temperatures can be estimated by several methods: (a) from the appearance of the UO₂ fuel (fuel grain growth and evidence of melting), (b) relating the temperatures at which phase changes occur in zircaloy cladding to the observed structures, (c) correlating the overall composition of the material examined, the number and appearance of the phases present, and the composition of the individual phases to existing phase diagrams, and (d) determining the presence of prior molten structural and control rod materials. The methods for determining peak temperatures are discussed in Sections 3.2.2.1 through 3.2.2.4, and the temperature estimates are discussed in Section 3.2.2.5.

3.2.2.1 <u>Fuel Grain Growth and Melting</u>. Fuel grain growth is highly time-temperature dependent and can be calculated using correlations of time at temperature for a given fuel burnup. These calculations are based on an integrated total time at a given temperature, however, and are difficult to use for transient conditions and multiple temperature excursions. Figure 17 shows the relationships between grain size and temperature for times at temperature of 20 and 40 min. The relationships shown in this figure are for initial grain size and burnup at the time of the accident, and for a range of times at temperatures that have been predicted for the accident. Therefore, an indication of what the temperatures were can be made by comparing and correlating measured grain size with Figure 17.

Other indications that fuel was exposed to high temperatures (>2000 K) are the rearrangement of pores and grain structure. As the

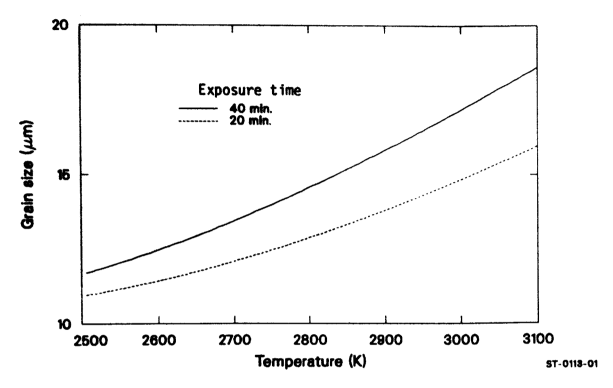


Figure 17. Plot showing the relationship between fuel pellet grain size and times at temperature.

melting point of the UO_2 fuel (3120 K for $\mathrm{UO}_{2.00}$) is approached, the fuel grains become almost pore-free as pores agglomerate at grain boundaries, sharp corners on the equiaxed grains become rounded, and the grains separate. When the fuel melts, the sintering porosity and grain boundaries disappear, leaving an amorphous densified material. With superheating above the melting point, bubbles of UO, UO₂, and UO₃ vapor form in the molten fuel--which creates a foamy appearance that survives cooldown to room temperatures. (Molten fuel foaming is aided by fission product gases and vapors in high burnup fuel.)

3.2.2.2 Zircaloy Phase Changes with Temperature. Zircaloy cladding progresses through several distinct microstructural changes with increasing temperature. The original (as-fabricated) zircaloy microstructure consists of alpha-zircaloy (α -Zr) grains elongated in the axial direction (referred to as a stress-relieved microstructure). This structure is observed with peak temperatures less than 920 K. With increasing temperature, the α -Zr grains recrystallize and become equiaxed (920 < T < 1105 K). Transformation from α -Zr to the β -Zr phase begins about 1105 K and ends about 1245 K. The temperature at which this transformation occurs increases with increasing oxygen content in the Zr. The β -Zr transforms back to α -Zr as temperatures are reduced below 1105 K; however, the microstructural appearance of the transformed β -Zr grains (referred to as "prior β -Zr") is distinctly different from the $\alpha\text{-}Zr$ grains that were not exposed to temperatures greater than 1105 K. The prior β -Zr structure appears as large, equiaxed grains with α -Zr platelets within each grain. The two-phase α + prior β -Zr mixture is observed in materials exposed to 1105 < T < 1245 K.

Several phenomena occur between 1245 < T < 2030 K. Interaction of the zircaloy cladding with steam on its outside surface and $\rm UO_2$ fuel on its inside surface becomes significant above 1245 K, and oxidation of the cladding occurs. Two layers are formed on the outside surface: first, a layer of α -Zr is formed that has been stabilized at high temperatures because of oxygen uptake; this layer is termed oxygen-stabilized alpha-zirconium [α -Zr(0)]; and then a layer of ZrO $_2$ is formed outside

the α -/r(0) layer. The longer the cladding is exposed to temperatures greater than 1245 K, the thicker these two layers can become. The entire wall thickness can eventually transform to ZrO_2 and α -/r(0). Frequently, there are two layers in the ZrO_2 phase. The inside layer [closest to the α -Zr(0)] contains a metallic α -Zr(0) phase. This indicates that the ZrO_2 was slightly hypostoichiometric at temperature and underwent eutectoid decomposition into stoichiometric $ZrO_{2.00} + \alpha$ -/r(0) during cooling. This decomposition only takes place if significant amounts (enough to be visible) of the ZrO_2 have been at temperatures above about 1760 K (Figure 18).

If fuel-cladding contact exists, interaction of the cladding with the $\rm UO_2$ fuel also occurs, resulting in oxidation of the cladding from the inside surface. The internal $\rm UO_2/Zr$ interaction results in the formation of the following reaction layer sequence:

$$[UO_2 + U] \rightarrow [\alpha-2r(0)_a + U,2r]$$

 $\rightarrow U,2r \ alloy \rightarrow \alpha-2r(0)_h$

where

- $a = the \alpha-Zr(0)$ next to the fuel
- b = the α -7r(0) near the center of the cladding.

These layers will then be followed by prior $\beta\text{-}Zr \to \alpha\text{-}Zr(0) \to ZrO_2$ or only $\alpha\text{-}Zr(0) \to ZrO_2$, depending on the amount of oxidation from the outside by the steam. The fuel is reduced by the Zr to form metallic $\alpha\text{-}Zr(0)$ and elemental uranium. ZrO_2 cannot form as a result of the $UO_2/7r$ reaction alone. The metallic uranium does not remain at the UO_2/α - $7r(0)_a$ interface, because it tends to interact with Zr low in oxygen and diffuses into the cladding to form a U,Zr alloy rich in uranium. The U,Zr alloy lies between two $\alpha\text{-}Zr(0)$ layers and is liquid

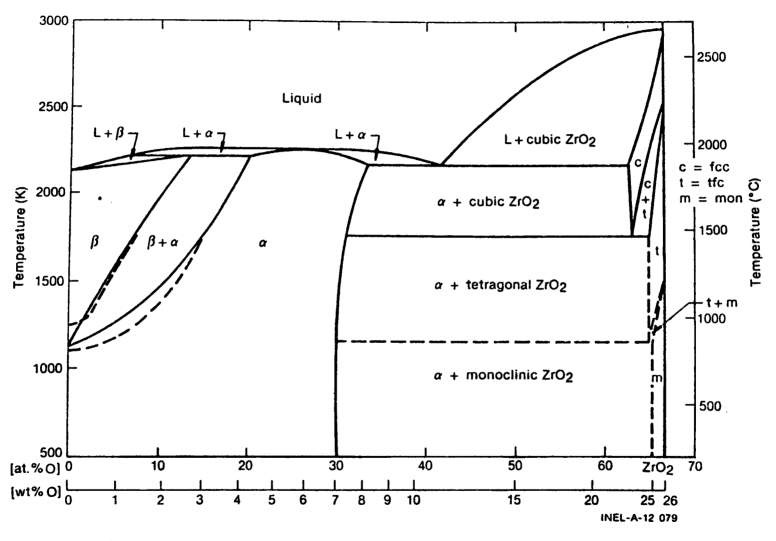


Figure 18. Zirconium-oxygen equilibrium pjhase diagram (Reference 10).

above about 1425 K, depending on the Zr content. This small quantity of liquid is a product of the solid state diffusion couple. The $\alpha\text{-}Zr(0)$ layer adjacent to the fuel contains small amounts of the U,Zr alloy, primarily along grain boundaries, but also as small globules within the $\alpha\text{-}Zr(0)_a$ grains. The $\alpha\text{-}Zr(0)_b$ layer adjacent to the prior $\beta\text{-}Zr$ contains no U,Zr alloy. During cooldown, the hypostoichiometric U0_2-x decomposes into stoichiometric U0_2.00 and additional metallic uranium. The two-phase $[\alpha\text{-}Zr(0)_a+\text{U,Zr}]$ layer is made up of relatively small, radially elongated $\alpha\text{-}Zr(0)$ grains compared to the $\alpha\text{-}Zr(0)_b$ layer [and $\alpha\text{-}Zr(0)_b$ in general] which is made up of large grains. 11-17

The boundary between the $[\alpha-Zr(0)_a+U,Zr]$ and U,Zr layers is distinct, but the interface between the U,Zr and $\alpha-Zr(0)_b$ layers can be very irregular. The formation of large U,Zr globules within the $\alpha-Zr(0)_b$ layers occurs only at temperatures greater than or equal to 1775 K; at lower temperatures, small spherical particles form. $^{12},13$

3.2.2.3 Interaction of $U0_2$ and Zircaloy. The $U0_2$ fuel and the zircaloy cladding comprise about 92% of the reactor core materials. Therefore, the interactions of these two materials with each other and with steam are the most important phenomenon to be considered in evaluating behavior of core materials. The zircaloy cladding can begin to melt at 2030 K, while stoichiometric UO, fuel does not melt until 3120 K; however, interaction of the Zr , UO_2 , and steam results in materials with intermediate melting temperatures. The melting point of Zr increases with increasing oxygen content; the melting temperature of oxygen-saturated a-Zr(0) is about 2245 K. A eutectic interaction takes place between oxygen-saturated α -Zr(0) and UO $_2$ at about 2170 K (27 at.% UO $_2$), and a monotectic interaction at about 2670 K (81 at.% UO $_2$). The UO $_2$ and ZrO₂ form a solid solution at high temperatures, and a 50/50 mole% composition melts at about 2809 K. Therefore, melting of the cladding and dissolution of the ${\tt UO}_2$ fuel (liquefaction) can begin at temperatures as low as 2030 K. Other low-temperature eutectics may form between the fuel rod materials (Zr and UO_2) and other core structural materials (Incone), stainless steel, and control rod materials).

The key to determining the peak temperatures reached and the possible scenario is the number, composition, and distribution of phases present in a prior molten material at room temperature. Neglecting effects of alloying elements, the observed phases should agree with the U-Zr-O ternary phase diagram. The ternary equilibrium phase diagrams of the U-Zr-O system at 1773 and 2273 K are shown in Figure 19. The equilibrium phases shown in the 1773 K diagram are representative of what would be seen at room temperature, because further phase changes at lower temperatures require very long times to reach equilibrium.

Interaction between UO $_2$ and Zr at 2273 K results in a single-phase homogeneous liquid or a two-phase U-Zr-O melt plus (U,Zr)O $_{2-x}$ solid (at >3.6-wt% oxygen). Upon cooling, the homogeneous melt will decompose into two metallic components: an α -Zr(O) phase containing some uranium and an U,Zr alloy containing little or no oxygen. 11,15,18

The heterogeneous melt also decomposes on cooldown, with the solid $(U,Zr)O_{2-x}$ particles having low Zr content decomposing into stoichiometric $(U,Zr)O_2$ and a U,Zr alloy, and the U-Zr-O melt decomposing into α -Zr(0) and U,Zr alloy.

The ongoing steam oxidation of the heterogeneous and homogeneous melts must be considered, as this affects the final microstructural appearance. If oxidation occurs after solidification, the α -Zr(0) will transform to ZrO₂ and the U,Zr alloy will transform to (U,Zr)O₂. Therefore, the homogeneous melt will transform from α -Zr(0) + U,Zr to ZrO₂ + (U,Zr)O₂, and the two phases should be distinguishable. The heterogeneous melt that oxidizes after solidification has an additional transformation of (U,Zr)O_{2-x} to (U,Zr)O₂.

The molten material also may absorb oxygen while in the liquid state. As the oxygen content of the melt increases, the temperature must increase for the material to remain molten, or a $(U,Zr)0_2$ solid will precipitate out. Complete oxidation of the melt at temperatures below 2800 K will result in a single-phase $(U,Zr)0_2$ solid solution. This material will no

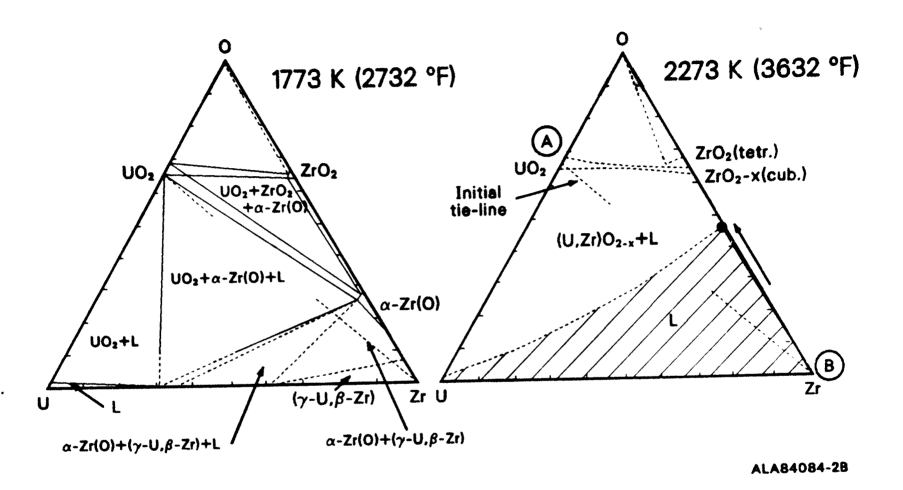


Figure 19. U-Zr-O ternary phase diagrams for temperatures of 1773 and 2273 K (Reference 20).

longer have the appearance of having been molten. Elongated grains will tend to form along the oxygen concentration and temperature gradients, and the material will generally contain few pores and be dense in appearance.

The ${\rm Zr0}_2$ and ${\rm U0}_2$ also can form a solid solution by diffusion at temperatures up to 2800 K and by mixing of molten oxides at higher temperatures. Figure 20 shows the ${\rm Zr0}_2$ – ${\rm U0}_2$ binary phase diagram. The minimum melting point of the solid solution occurs at about the 50/50 mole% mixture. Only one phase is apparent on cooling of the $({\rm U,Zr}){\rm O}_2$ mixture. If this material has been molten, it will generally show structures such as large pores and irregular or nonexistent grain structure.

The amount of interaction with other materials is another indication of whether the post-transient $(U,Zr)0_2$ solid solution was originally a metallic melt. If the metallic melt contacts a ceramic material $(U0_2)$ or $Zr0_2$, dissolution of the ceramic by reduction should occur. A ceramic melt contacting a ceramic solid will show very little interaction, although mixing can occur if the solid melts due to conducted heat.

3.2.2.4 Melting Temperature of Other Reactor Materials. Although the UO₂ fuel and zircaloy cladding comprise the major portion of the reactor core, the other reactor materials could play an important roll in fuel behavior. This is because the other reactor components have lower melting temperatures than the UO₂ fuel and many of these components can form eutectics with the fuel rod materials, thus, lowering fuel melting temperatures and changing physical properties. Table 8 presents the core material compositions along with the melting temperatures of the major constituents. The presence of prior molten control rod and structural materials in a particle would indicate peak temperatures of at least the melting point of that material. Unfortunately, phase diagrams do not exist for all possible combinations of the various reactor materials, and the effect of adding trace amounts of alloying elements on melting temperatures is not always quantified.

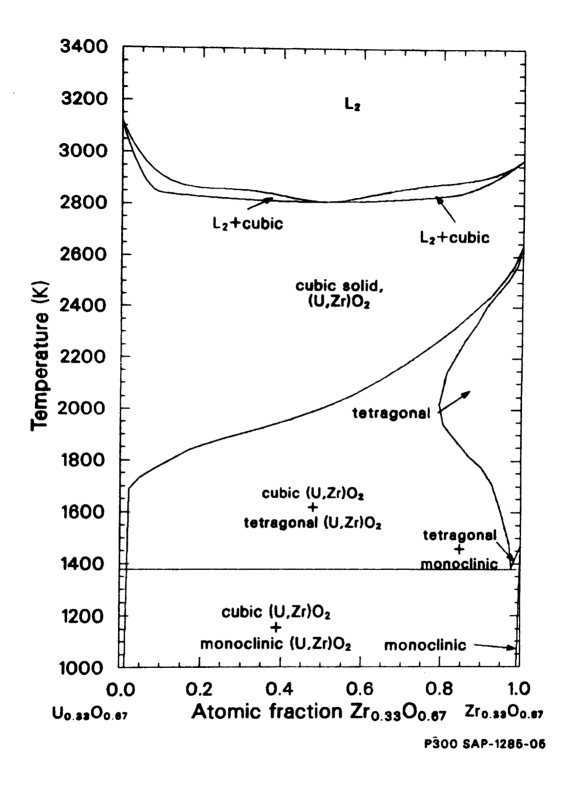


Figure 20. U02-Zr02 binary phase diagram (soludus and liquidus curves represent preliminary data from Reference 19).

TABLE 8. CORE MATERIAL COMPOSITION

	Material					Material					
	Wel	ght	_ Melt	11. 1 . 1.		<u>We1gl</u>	<u>1t</u>	Melt lemperature		Weight	
	<u>(kg)</u>	(%)	Temperature (K)	<u>Element</u>	Weight <u>Percent</u>		(kg)	(%)	(K)	<u>Element</u>	Percen
uo ₂	93050	74.0	3120	235 _Ս 238 _Ս 0	2.265 85.882 11.853	Incone 1-718	1211	1.0	1720	N1 Cr Fe Nb Mo	51.900 19.000 18.000 5.55 3.00
?trcaloy-4	23029	18.3	2030	Zr Sn Fe Cr O C N Hf S Al T1 V Mn N1 Cu	97.907 1.60 0.225 0.125 0.095 0.0120 0.0080 0.0078 0.0035 0.0024 0.0020 0.0020 0.0020	2r0 ₂	331	0.3	2960	Al Co Sl Mn N Cu C S	0.80 0.60 0.47 0.20 0.20 0.13 0.10 0.04 0.00
				W H Co B Cd	0.0020 0.0013 0.0010 0.000033 0.000025	Ag-In-Cd	2149	2.2	1050	Ag In Cd	80.0 15.0 5.0
Type 304 and indentified stainless steel	4636	3.7	1720	Fe Cr N1	0.000020 68.635 19.000 9.000	8 ₄ C-A1 ₂ O ₃	626	0.5	2320	A 1 O B C	34.3 30.5 27.5 21.5
•				Mn S1 N C Co P	2.000 1.000 0.130 0.080 0.080 0.045	84t Gd ₂ 03-U02.	131.5	0.1	2620 2400	6d U 0	10.2 71.7 12.0
				P S		Gd ₂ O ₃			26/0		

3.2.2.5 <u>Temperature Estimate Results</u>. The estimated peak temperatures for the 29 particles are listed in Table 7, along with other relevant data about each particle. These estimated temperatures are presented again in lable 9, along with comments about key indicators used for estimating the temperatures. (for more detail see Appendix C.) These estimates are for the highest temperatures on the particle and may not represent those for the bulk of the particle. Only three of the particles examined do not contain molten Zr or Zr, U, 0 mixture. One of these is a prior molten Zr particle, and the other two are sections of cladding oxidized to α -Zr(0) but not melted; therefore, phase changes occurring at low temperatures (<2030 K) in Zr could not be used as indicators.

Many of the particles examined contain UO₂ fuel, and usually the grain growth is minimal (maximum measured was 29 µm, with the original grain size being 10 µm). This indicates that much of the fuel probably remained at fairly low temperatures (<2000 K) during the accident or was exposed to high temperatures for only a short time. However, most of these particles also have areas of prior molten material, indicating large temperature gradients over very small distances. (The particles all are less than 4 mm in diameter.) The molten material relocated to a large extent and interacted both chemically and mechanically with the cooler materials.

The reader is reminded that the particles selected for detailed metallurgical examination were chosen either because they were representative of a basic particle type or because they were unique. They are not a statistical representation of the number of particles estimated to have experienced a given peak temperature. The particle type distribution shown in Table 6 (see Section 3.1.5) represents all of the larger particles in ten of the core debris grab samples and is more representative of the core debris. Based on this data and data from the detailed metallurgical examinations, it is estimated that about 80% of the debris material remained at fairly low temperature (<2000 K) due to a lack of restructuring, with the other 20% experiencing temperatures up to fuel melting.

TABLE 9. PEAK TEMPERATURE ESTIMATES FOR PARTICLES FROM THE TMI-2 CORE DEBRIS GRAB SAMPLES

<u>Particle</u>	Temperature Range ^a (K)	Comments
1A	2170~2245	No melting of $U0_2$; ballooned α -Zr(0); eutectic Zr-U interaction.
18	2170-2600	No melting of UO ₂ ; slight grain growth; eutectic Zr-U interaction.
1E	2810-2960	No melting of ZrO_2 ; melting of α - $Zr(0)$ cladding; prior molten (U,Zr) O_2 .
Н	>2810	Dissolution of $(U,Zr)O_2$ by porous heterogeneous $U-Zr-O$ melt.
3L	>2850	Above the liquidus temperature for $U-Zr-O$ ceramic.
3M	>2850	Areas of (U,Zr)O ₂ melt; contains trace Cr, Ni, Fe.
4A	~2810	Foamy U-Zr-O molten ceramics contacting UO ₂ ; may be slightly lower due to impurities.
4 B	~2810	Foamy U-Zr-O molten ceramic contacting UO ₂ fuel; may be slightly lower due to impurities.
40	>2850	Molten (U,Zr)O ₂ ; dense.
5E	~2810	Foamy U-Zr-O molten ceramic contacting fuel.
6B	<2245	No melting of α -Zr(0).
6C	2170-2960	No melting of ZrO2; UO2-Al2O3 eutectic interaction.

TABLE 9. (continued)

<u>Particle</u>	Temperature Range ^a (K)	Comments
60	2170-2810	No fuel melting; metallic U-Zr-0 melt, oxidized to $(U,Zr)0_2$ by reducing oxidized fuel to $U0_2$.
6E	~2000	Oxidized cladding fragment; no melting.
6F	2170-2810	Porous heterogeneous U-Zr-O melt attached to $(U,Zr)O_2$, ferromagnetic ingots.
7A	~2810	Foamy U-Zr-O molten ceramic; may be lower due to impurities.
78	2170-2245	No melting of UO_2 or α -Zr(0); eutectic U-Zr interaction.
7E	~2810	Foamy U-Zr-O molten ceramic
8A	~2810	Foamy U-Zr-O molten ceramic.
8C	>2170	Oxidized U-Zr-O melt.
8E	~2810	Foamy U-Zr-O melt contacted later by metallic Zr melt.
8н	>2850	Areas of dense (U,Zr)O ₂ melt containing trace Fe.
90	~28 10	Foamy U-Zr-O ceramic melt.
96	>1233	Molten Ag.
10A	~3100	Areas of dense (U,Zr) 0_2 melt with 5% Zr 0_2 .
108	>2810	U-Zr-O ceramic melt on UO_2 fuel pellet.
10F	>2850	Regions of dense and foamy U-Zr-O melt.

TABLE 9. (continued)

<u>Particle</u>	Tem	perature Range ^a (K)	Comments				
118	>2810		Foamy U-Zr-O melt.				
110		2170-3100	No melting of UO2; small region of prior molten (U, Zr) O_X .				
a. Explanation of temperature ranges:							
T ₁ -T ₂	:	peak temperature greater than T_1 but less than T_2 (i.e., some feature indicated temperature greater than T_1 and some feature indicated temperature less than T_2)					
~11	:	peak temperature	about T ₁				
>T ₁	:	peak temperature	at least T ₁				
<t<sub>1</t<sub>	:	Peak temperature	less than T ₁				

The key indicators used to estimate peak temperatures in the ranges of interest are shown in Figure 21. Most of the temperature estimates range between 2030 and 3120 K. The specific microstructures recognizable for materials exposed to temperatures in this range were discussed in Section 3.2.2.3. With some overlap, the particles examined fall into the following categories:

- Some particles show a prior molten, metallic U-Zr-O mixture in contact with a UO_{2} fuel pellet fragment. Therefore, the melt temperature before contact had to have been greater than 2170 K. the α -Zr(0) eutectic temperature. In some cases, an upper melt temperature limit can be determined, depending on behavior of other particle components. If a substantial metallic melt has flowed between ballooned cladding and fuel without melting the a-Zr(0) cladding, then the maximum melt temperature was less than 2245 K (Figure 22). Similarly, if a large metallic melt has surrounded a UO, fragment (dissolving a portion by eutectic decomposition) without melting any pure UO, by heat conduction, the maximum melt temperature would have been less than 3120 K. However, in many instances, the melt size is much smaller than the substrate material, in which case the large heat sink chilled the melt very quickly and no upper temperature limit can be established.
- One particle (Particle 6C, Figure 23) shows evidence of interaction with the Al_20_3 poison rod material. A eutectic can form between $u0_2-zr0_2-Al_20_3$ at a temperature as low as 2170 K. This particle contains $zr0_2$ which had not melted, indicating a temperature range of 2170 to 2960 K.
- Several particles show evidence of prior molten $(U,Zr)0_2$ ceramic, for which a minimum melting temperature of 2810 K has been established at KfK in the Federal Republic of Germany. At KfK, the melting temperature was found to increase only for compositions near $U0_2$ (melts at 3120 K) or $Zr0_2$ (melts at

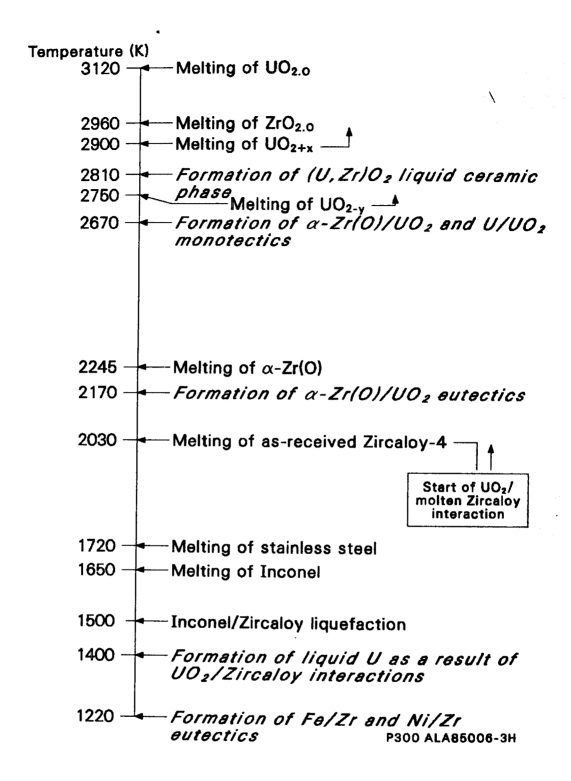


Figure 21. Chemical interactions and formations of liquid phases in the $UO_2\text{-}Zr$ system with increasing temperature.

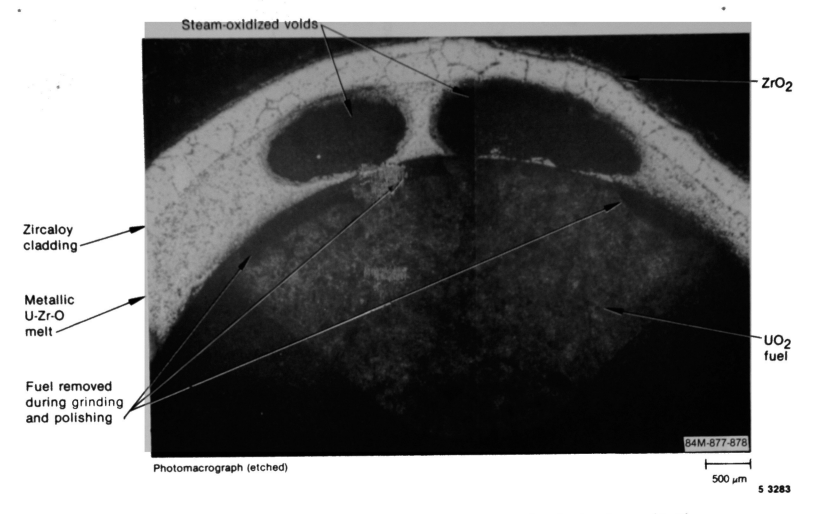


Figure 22. Photomacrograph of Particle 1A (H8, surface) showing melt flow between UO₂ fuel and ballooned cladding.

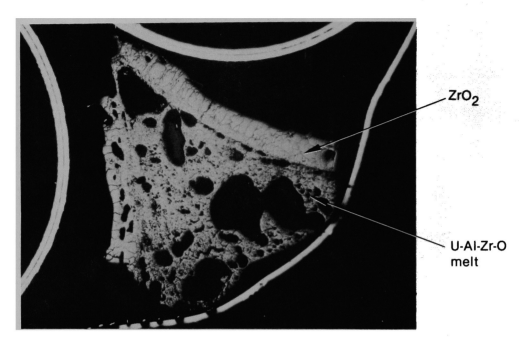


Figure 23. Photomacrograph of Particle 6C (E9, 56 cm) showing interaction of fuel rod materials with ${\rm Al}_{2}{\rm O}_{3}$.

- 2973 K). This work also demonstrated that a foamy structure with large voids is formed very near the melting point, but the $(U,Zr)O_2$ displays a much more dense structure upon exposure to temperatures above the liquidus (see figure 20). Two of the particles in this category show dendritic freezing in the $(U,Zr)O_2$ --separation of the UO_2 - ZrO_2 homogeneous liquid into U-rich and Zr-rich phases--indicating slow cooling from above the liquidus. Examples of the foamy $(U,Zr)O_2$ structure, the dense, mixed ceramic melt, and the dendritic freezing are presented in Figures 24, 25, and 26, respectively.
- Particle 10A (Figure 27) also contains regions of prior molten 0 (U,Zr)0₂, but a large part of this material contains slightly less than 2 atom% Zr (which corresponds to about 5-mole% ZrO₂). The small amount of Zr or other impurities indicates a liquidus or solidus temperature approaching UO2 melting temperatures (2750 to 3120 K, depending on 0 to U ratio, See Figure 31). There is a small central region of UO, that has been restructured but not melted. This particle appears to have been a piece of fuel that interacted with ZrO, near the UO, melting point. The prior molten ceramic material has a dense, almost glassy appearance, indicating temperatures somewhat above the liquidus temperature and then rapid cooling. No grain structure could be found outside of the central restructured zone, either during the optical examination using etches or during the SEM examination.
- Other particles contain considerable porosity that is not related to the foamy structure just discussed. As shown in Figure 28, the α -Zr(0)/U0₂ isopleth, when α -Zr(0) has dissolved appreciable amounts of U0₂ and when the melt is above 2673 K, the molten mixture is composed of two partially miscible liquids. Between 2673 and 2173 K, one liquid (L1) transforms to (U,Zr)0_{2-X} solids, forming a slurry, while below 2173 K, the

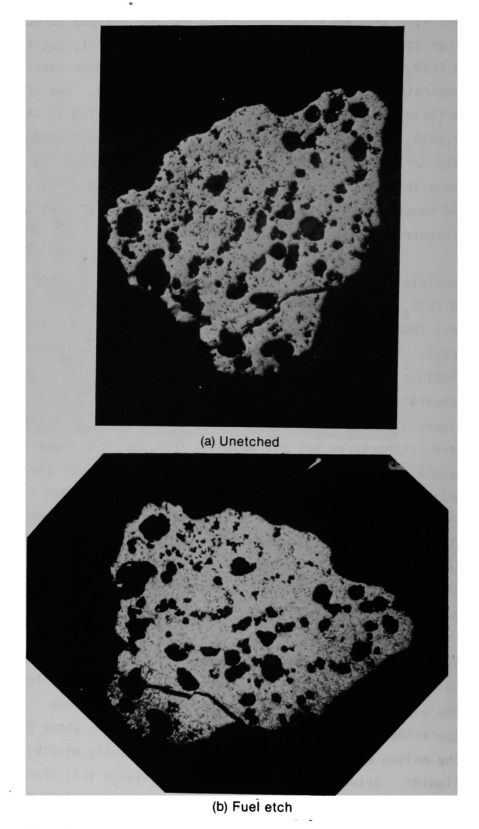


Figure 24. Photomacrograph of Particle 7A (H8, 36 cm) showing foamy U,Zr,O melt.

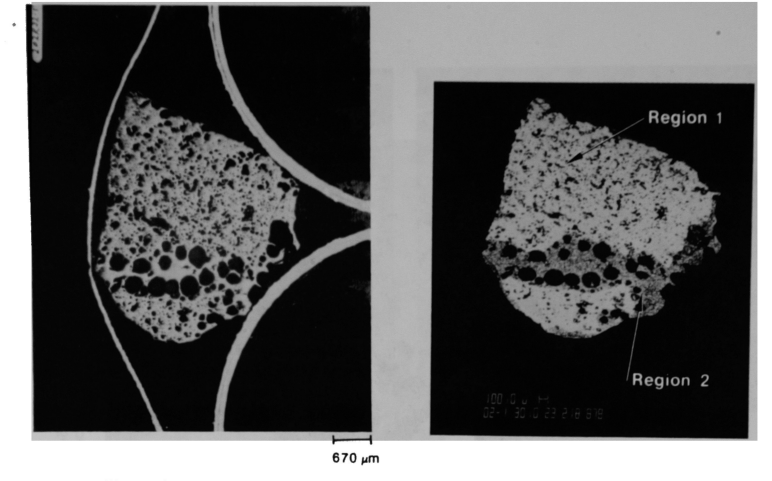


Figure 25. Photographs of Particle 10F (E9, 74 cm) showing the presence of of two distinct regions: Region 1 is dense U,Zr,O melt and Region 2 is foamy U-Zr-O melt.

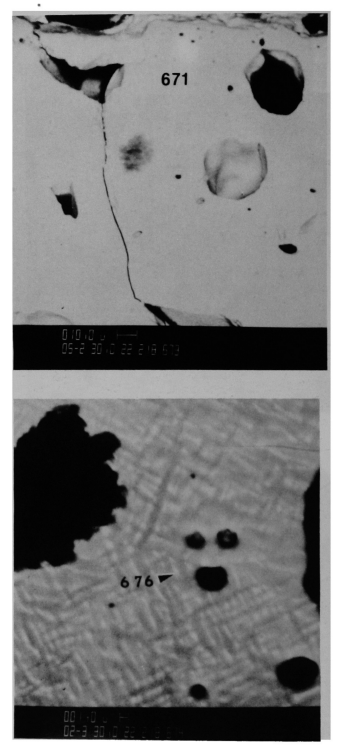


Figure 26. SEM back scattered electron image of Particle 3L (H8, 56 cm) showing dendritic freezing.

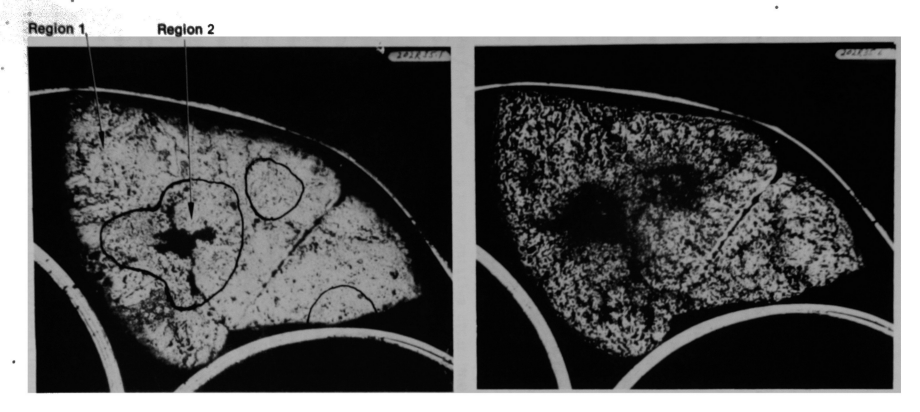


Figure 27. Photomicroggraphs of Particle 10A (E9, 74 cm) showing prior molten material in Region 1. Region 2 is restructured, but not melted, material UO₂.

(b) Etched

(a) Unetched

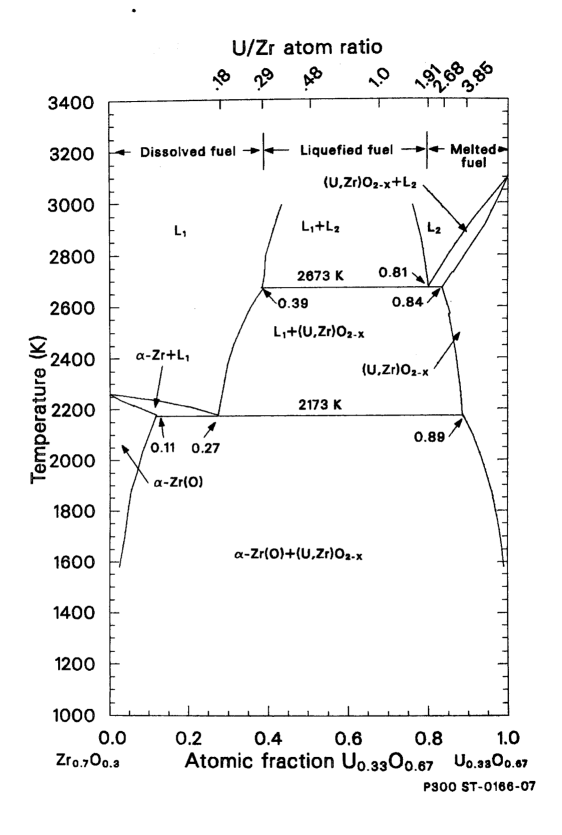


Figure 28. Quasi-binary α -Zr(0)/U02 phase diagram (Reference 20).

other liquid (L1) converts to α -Zr(0). Both the liquid mixture and the slurry display a strong tendency to chemically attach to UO_2 , $(U,Zr)O_2$, and other solids.

figure 28 also indicates that U-to-Zr atom ratios are a valuable indicator of melt temperatures, especially since oxygen concentrations were probably changed; by subsequent steam oxidation. This wetting property is so dominant that, upon cooldown or quenching, shrinkage voids are created as the heterogeneous melt is pulled apart. Bubbles of gases and metal vapors also may contribute to void formation, though these pore sources have not yet been positively confirmed. In any case, heterogeneous melt porosity is typically irregular in configuration, with elongations, jagged edges, and other features unlike the smooth spherical pores inside foamy material. One example of a porous, heterogeneous melt is shown at the base of Particle 6F in Figure 29.

Particle 6F also is representative of another broad category of particles that suggest occurrence of either multiple heatups during the TMI-2 transient or prolonged candling of molten material down the outsides of fuel rods, with intermittent solidification and oxidation of relocated droplets. The upper crown of Particle 6F is single-phase (U,Zr)0, that was formed either by ceramic melting or oxidation of a metallic U-Zr-O In either case, the (U,Zr)0, solidified before being coated by the porous heterogeneous melt. The porous melt, in turn, solidified by cooling before being contacted by a Ni-rich metallic melt that created the three ferromagnetic ingots. On other particles, melt penetration is frequently observed along grain boundaries of previously solidified material. multiple temperature ramps and a lengthy candling sequence are plausible explanations, but this particle category definitely establishes that fuel liquefaction, relocation, and solidification did not occur within a single brief interval.

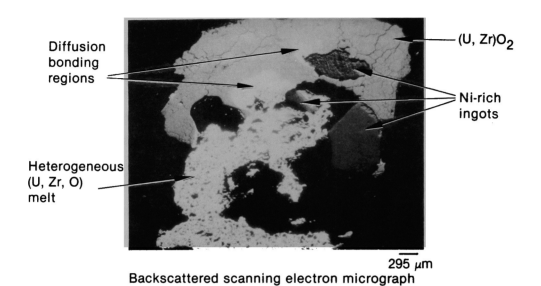


Figure 29.

Particle 6F (E9, 56 cm) shows where solidified (U, Zr)0 $_2$ was contacted by a porous melt that solidified before contact by a Ni-rich melt.

- As mentioned above, (U,Zr)O₂ regions can form by oxidation of metallic melts. Particle 6D shows an example of such material (Figure 30). A small amount of metallic melt contacted a large fuel pellet fragment, reducing the adjacent hyperstoichiometric fuel while the melt oxidized to single-phase (U,Zr)O₂. Note that (U,Zr)O₂ formed by this liquid-state oxidation mechanism is very difficult to distinguish from a rapidly cooled, superheated ceramic melt, unless the melt has coated fuel or ZrO₂. Whereas a metallic melt must reduce the adjacent oxide, a ceramic melt could not alter the oxygen concentration of the coated substrate.
- oxidized fuel. As detailed in Appendix C, a second fuel phase was exposed by etching this particle. Quantitative Scanning Auger Spectroscopy (SAS) was used to measure an average oxygen concentration in the fuel matrix of 71 at.%, which converts to UO_{2.44}--well above stoichiometric UO₂. According to the U-O phase diagram in Figure 31, the two phases are U₄O₉ and UO_{2.6} (either U₅O₁₃ or U₈O₂₁). Therefore, oxygen gas or hydroxide radicals must have diffused into the UO₂ above 1900 K. Additional SAS measurements confirmed that fuel next to the adherent melt was reduced back to UO₂ while the melt became (U,Zr)O₂, which agrees with the results of the metallography.

Prior molten structural materials were observed in a few samples, defining the lower temperature range for those particles. An upper temperature could not be established for particles with no unmelted portions. Also, the temperatures quoted are the highest peak temperature for the materials that make up a given particle. Thus, the peak temperature quoted could be for a small layer of material adhering to the surface of an otherwise low temperature particle.

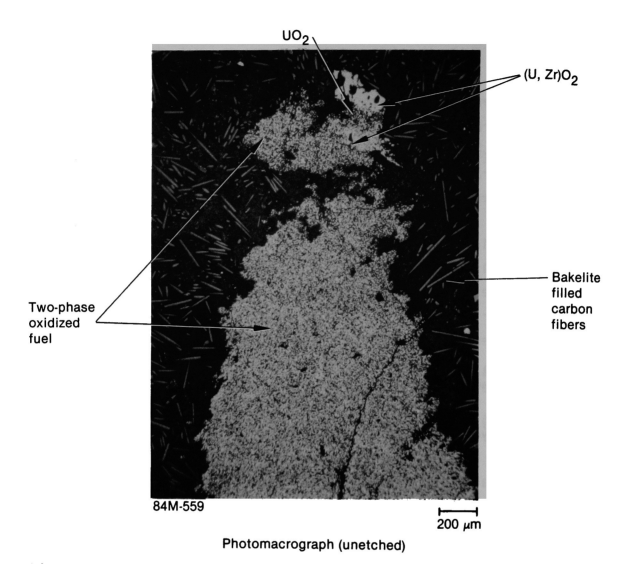


Figure 30. Particle 6D (E9, 56 cm) shows where the oxidized fuel mixture of U_4O_9 and U_5O_{13} have been reduced to UO_2 at one corner by contact with an initially metallic U-Zr-O melt.

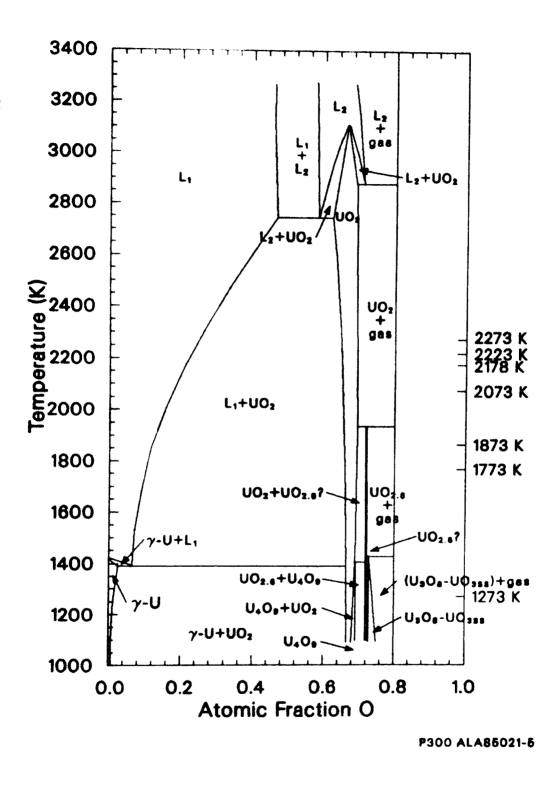


Figure 31. Uranium-oxygen phase diagram (Reference 21).

3.2.3 Interactions of Fuel Rod and Non-Fuel Rod Materials

The fuel rods, including $\rm UO_2$ pellets and zircaloy cladding, made up about 92% of the original TMI-2 core inventory. The remaining 8% was structural and control material. Table 10 lists the core materials by elemental weight percent, excluding oxygen.

Most of the particles contain at least some traces of structural materials (Cr, Fe, and Ni), and in some cases structural materials are a significant part of the particle (see Particles 4A, 6C, 7A, or 8C in Appendix C). These materials are commonly found mixed intimately with prior molten fuel rod materials, primarily at grain boundaries or in voids. However, it was also common to find traces of Fe in the base material (see Table 7 for a listing of particles that contain Fe in the base material). The almost universal occurrence of structural material with prior molten fuel rod materials indicates that strong reactions took place between fuel and structural materials. These reactions have been observed by Hagen et al. in experiments with electrically heated fuel rods and Inconel grid spacers. 22

Structural materials (Ni with small amounts of Fe) also were found in non-oxidized metallic form, mechanically bonded to fuel rod materials, as seen in Particle 11B. Ni alloyed with small amounts of Sn and occasionally fission products (Ru and Tc) was found scattered throughout many of the particles in small, round, metallic (non-oxidized) inclusions. It is possible for Ni to be separated from the other Inconel or stainless steel constituents through a membrane separation process, whereby the Fe and Cr preferentially diffuse into the solid (U,Zr)O₂. An example of this can be seen in Particle 7E, along an edge that contains many small regions of Ni surrounded by material high in Cr and Fe.

Control rod materials (Ag, In, and Cd) were not as commonly found as the structural materials. Ag was occasionally found, primarily in inclusions, and there were two minor occurrences of In and Cd. However, major portions of three particles (Particles 8H, 9D, and 9E) were metallic

TABLE 10. TMI-2 CORE COMPOSITION BY ELEMENTAL WEIGHT PERCENT

<u>Element</u>	Weight Percent ^a
Ų	71.77
Zr	19.89
Fe	3.01
Ag	1.75
Cr	1.00
Ni	0.91
In	0.33
Sn	0.32
Α٦	0.18
В	0.14
Cd	0.12
Mn	0.09
Nb	0.05
St	0.04
С	0.04
Mo	0.03
Gd	0.01
Ti	0.01
N	0.01
Co	0.01
S	<0.01
Ca	<0.01
P	<0.01
Hf	<0.01
٧	<0.01
W	<0.01

a. The initial oxygen content (9.02 wt.% total in fuel, poison rods, ZrO₂ spacer pellets) was omitted in calculating these weight percents to avoid confusion with the oxidation that occurred during the accident.

Ag containing Ni-Sn inclusions. Particle 9G was composed completely of the Ag material, and Particles 8H and 9D contained ceramic fuel rod material mechanically bonded to the Ag material. This lack of mixing with fuel rod materials is probably due to the early failure of the control rods, while the rest of the fuel rod bundle was still intact. In addition, eutectic interactions were probably inhibited by oxide layers.

The major source of Al in the core was from alumina-boron carbide $(B_4C-Al_2O_3)$ burnable poison rods. Al also is a minor alloying element in Inconel, and there were other minor sources. However, Al was found with other structural materials in over half of the particles that contained structural materials. If the source of Al in these particles was Inconel, then there would have had to have been a segregation process at work concentrating the Al. However, if the source of Al was the poison rods, then there first would have had to have been eutectic interactions between Inconel grid spacers, zircaloy cladding, and alumina, causing liquefaction and flow, and then eventual interaction with fuel rods. Particle 6C contains exceptionally large amounts of Al, indicating a possible direct interaction with a poison rod. There are areas on this particle where positive elemental identification is not possible because of nonconductive properties, but there are indications that these areas are alumina.

3.3 Chemical Analyses

Elemental analysis was performed on the recombined bulk samples and particles and aliquots using inductively coupled plasma spectroscopy (ICP). The original elemental composition and total weights of principal components for the core and a control rod fuel assembly are presented in Table 11. The elements for which analysis was performed by ICP also are identified. The elements were selected for analysis to characterize the five groups of materials present in the core; uranium fuel and zircaloy cladding, Ag-In-Cd control rod materials, burnable poison rod materials(Al, Gd, and B), structural materials (stainless steel and Inconel), and Te.

TABLE 11. TMI-2 REACTOR CORE AND CONTROL ROD FUEL ASSEMBLY COMPOSITION

					
Material		Weight			Weight
(Weight)	Element	Percent	<u>Material</u>	<u>Element</u>	Percent
, UO ₂	U-235ª	2.265	Inconel-718	Nia	51.900
(93,050 kg)	U-238a	85.882	(1,211 kg)	Cra	19.000
(526 kg) ^b	0	11.853	$(6.8 \text{ kg})^{5}$	Fea	18.000
(020 113)	•		(Nb	5.553
				Мо	3.000
Zircaloy-4	Zra	97 .907		Ti	0.800
(23,029 kg)	Sna	1.60		ΑΊa	0.600
(125 kg) ^b	Fea	0.225		Co	0.470
	Cra	0.125		Sta	0.200
	0	0.095		Mna	0.200
	C	0.0120		N Guð	0.130
	N	0.0080		Cua	0.100 0.040
	Hf	0.0078		C S	0.040
	S	0.0035		3	0.007
	Alg	0.0024			
	Ţi	0.0020 0.0020	ZrO2	Zra	74.0
	Mug A	0.0020	(331 kg)	Ü	26.0
	Nig	0.0020	(0 kg)	Ū	2010
	Cua	0.0020	(O K9)		
	W	0.0020			
	Ĥ	0.0013	Ag-In-Cd	Aga	80.0
	Co	0.0010	(2.749 kg)	Ina	15.0
	Bā	0.000033	$(43.6 \text{ kg})^{b}$	Cda	5.0
	Cda	0.000025			
	Ua	0.000020			24.22
			B4C-A1203	Ala	34.33
			(626 kg)	0	30.53
	- 3		(0 kg) ^b	B C	27.50 7.64
Type 304	Fea	68.635		C	7.04
stainless	Cra	19.000			
s tee 1	Nia	9.000	Cd=0=-110=	Gda	10.27
(676 kg) and	Mna	2.000	Gd ₂ O ₃ -UO ₂	U	77 .7 2
Unidentified	Sta	1.000 0.130	(131.5 kg) (0 kg) ^b	ŏ	12.01
stainless	N	0.130	(0 /9)	· ·	
steel	C	0.080			
(3,960 kg)	Co P	0.045			
(16.8 kg) ^b	Š	0.030			
	•				

a. Elements for which ICP analysis was performed.

b. Weight of material in a control rod fuel assembly.

The objectives of the elemental analysis include (a) characterizing the homogeneity of the core debris bed, both axially (0 through 94 cm in depth) and radially [from the H8 (center) to the E9 (mid-radius) locations—a distance of ~70 cm], (b) determining the retention or release of highly volatile core materials (e.g., Ag-In-Cd), and (c) characterizing the relocation of core materials and Te.

Results of the chemical analysis as they relate to the present condition of the core and the accident scenario, are evaluated in Section 4.

3.3.1 Summary of Examinations

Results of the ICP elemental analysis for the recombined bulk samples are presented in Tables 12 and 13. Results are presented in weight percent (wt%) of the dissolved material. Insoluble material weights are listed in Table E-12 (Appendix E). The insoluble material was less than 3% for any sample. The weight percents when added together do not equal 100%, because other elements expected to be present (e.g., oxygen) were not measurable by this method. The principal elemental components identified are U and Zr, with smaller amounts (generally <5 wt%) of other elements. The total uncertainty associated with the results of the elemental analysis of each recombined bulk sample is 10 to 15%. Factors contributing to the total uncertainty include (a) the presence of insoluble material, (b) ICP analysis uncertainty, (c) uncertainties associated with sample weight and volume measurements, and (d) presence of unmeasurable sample components.

Results of the elemental analysis of the particles and aliquots are contained Appendix D. The elements measured are the same as those shown in Tables 12 and 13 for the recombined bulk samples. The total uncertainty associated with the analysis of each particle and aliquot is generally 10 to 15%, because of the factors listed above. However, as noted in Appendix D, the total uncertainty for some particles and aliquots ranges from 30 to 50% because of the small (<10 mg) portions examined or sample losses incurred during the dissolution and analysis. Sample loss was

TABLE 12. ELEMENTAL CONTENT OF THE RECOMBINED BULK SAMPLES FROM THE H8

CORE LOCATION (wt%)

Element	Sample 1 $(23.56 \text{ g})^{\text{b}}$	Sample 7 (44.62 g) ^b	Sample 3 (50.10 g) ^b	Samples 8 and 9 ^C
Ag Al B Cd Cr Cu	1.6 E-1 3.37 8.02 E-1 <5.9 E-3 3.9 E-1 <1.5 E-2	1.9 E-1 <4.6 E-2d 2.8 E-1 <2.3 E-3 3.7 E-1 5.7 E-3	1.5 E-1 <4.8 E-2 9.3 E-2 <2.4 E-3 1.2 E-1 1.2 E-2	1.6 E-1 <6.0 E-2 1.17 <3.0 E-3 2.9 E-1 7.6 E-3
Fe Gd In Mn Mo Ni	1.19 8.2 E-2 <1.8 E-1 4.1 E-2 <5.9 E-2	1.17 5.7 E-2 <6.9 E-2 3.7 E-2 <2.3 E-2	5.0 E-1 5.8 E-2 <7.2 E-2 1.9 E-2 <2.4 E-2	9.9 E-1 5.7 E-2 6.6 E-2 3.4 E-2 <3.0 E-2
Nb Sie Sn Te U Zr	1.38 <1.8 E-1 3.84 <2.9 E-1 <1.2 E-1 7.03 E+1 1.59 E+1	1.6 E-1 3.63 E-1 <1.1 E-1 <4.6 E-2 7.39 E+1 1.76 E+1	1.7 E-1 1.7 E-1 1.39 <1.2 E-1 <4.8 E-2 7.25 E+1 8.24	3.3 E-1 1.7 E-1 2.8 E-1 <1.1 E-1 <4.6 E-2 7.23 E+1 1.42 E+1

a. In order of depth into the debris bed.

b. Weight of sample material dissolved.

c. Samples 8 and 9 were combined, as they had been mixed during dissolution.

d. Less than (<) values are the lower limit of detection for the sample analyzed.

e. Silicon contamination from the glassware occurred during the dissolution process.

TABLE 13. ELEMENTAL CONTENT OF THE RECOMBINED BULK SAMPLES FROM THE E9 CORE LOCATION^a (wt%)

Element	Sample 4 $(5.11 \text{ g})^{\text{b}}$	Sample 5 (16.64 g) ^b	Sample 6 (45.97 g) ^b	Sample 10 (57.44 g) ^b	Sample 11 (42.54 g) ^b
Ag	1.5 E-1	5.4 E-2	1.4 E-1	1.3 E-1	1.9 E-1
Al	1.1	<6.7 E-2 ^c	<7.7 E-2	<2.8 E-2	<7.2 E-2
B	3.00	1.4 E-1	1.6 E-1	1.70 E-1	<5.4 E-2
Cd	<2.1 E-2	<3.4 E-3	<3.9 E-3	<1.4 E-3	<3.6 E-3
Cr	<6.3 E-2	9.4 E-2	1.0 E-1	1.45 E-1	2.1 E-1
Cu	<5.3 E-2	<8.4 E-3	<9.7 E-3	5.6 E-3	<8.9 E-3
Fe	5.1 E-1	2.1 E-1	3.3 E-1	4.33 E-1	8.3 E-1
Gd	<1.5 E-1	4.7 E-2	5.4 E-2	5.6 E-2	<2.5 E-2
In	<6.3 E-1	<1.0 E-1	<1.2 E-1	<4.3 E-2	<1.1 E-1
Mn	2.1 E-1	1.4 E-2	1.6 E-2	1.4 E-2	2.1 E-2
Mo	<2.1 E-2	<3.4 E-2	<3.9 E-2	<1.4 E-2	<3.6 E-2
Ni	<1.1 E-1	2.7 E-1	2.4 E-1	2.55 E-1	3.9 E-1
Nb	<6.3 E-1	<1.0 E-1	2.7 E-1	9.8 E-2	<1.1 E-1
Sid	2.20 E+1	5.0 E-2	1.36	2.49 E-1	3.9 E-1
Sn	<1.1	<1.7 E-1	<1.9 E-1	1.4 E-1	<1.8 E-1
Te	<4.2 E-1	<6.7 E-2	<7.7 E-2	<2.8 E-2	<7.2 E-2
U	8.32 E+1	8.31 E+1	7.08 E+1	7.34 E+1	7.76 E+1
Zr	1.14 E+1	4.31	8.25	7.19	9.69

a. In order of depth into the debris bed.

b. Weight of sample material dissolved.

c. Less than (<) values are the lower limit of detection for the sample analyzed.

 $[\]mbox{\bf d.}$ Silicon contamination from the glassware occurred during the dissolution process.

determined by comparing the elemental U content obtained by ICP with results of the fissile/fertile material (235 U + 238 U) analysis performed on the intact particle or aliquot (Appendix E, Table E-11).

X-ray diffraction analysis was performed on some particles; however, due to problems associated with the technique (detailed in Appendix A), results were of limited value and, therefore, are not reported.

3.3.2 Chemical Composition of the Core Debris

Although individual particles of the core debris are quite heterogeneous on a microstructural scale, as discussed in Section 3.2, the composition of the debris bed is principally a mixture of U and Zr, with lesser amounts of other elements, similar to that listed in Table 11. The average elemental concentrations measured for the recombined bulk samples were used to characterize the homogeneity of the debris bed; results of the particle and aliquot analysis provide indications of the homogeneity of individual particles, or aliquots. Retention and release of volatile components (e.g., Ag-In-Cd) was estimated by comparing measured elemental concentrations with their original concentrations in the fuel assembly (see Table 11). Evidence of relocation of core components was determined by the wide distribution of less abundant constituents (e.g., Gd) in the debris bed.

Evaluation of the elemental analysis results is divided into characterization of the elemental constituents of the five principal groups of materials present in the core: (a) uranium fuel and zircaloy cladding, (b) Ag-In-Cd control rod materials, (c) burnable poison rod materials ($B_4C-Al_2O_3$ and $Gd_2O_3-UO_2$), (d) structural materials (stainless steel and Inconel), and (e) tellurium (the elemental analysis is used to evaluate the behavior of fission product Te).

3.3.2.1 <u>Uranium and Zirconium</u>. The reactor core is composed principally of U (82,000 kg) and Zr (22,500 kg). These elements account for approximately 84% of the total core weight. Table 14 lists the U and

TABLE 14. URANIUM AND ZIRCONIUM CONCENTRATIONS AND ZR/U RATIOS FOR THE RECOMBINED BULK SAMPLES

Sample	Depth into Debris Bed (cm)	Uranium Concentration ^a (wt%)	Zirconium Concentration ^b (wt%)	Zr/U Ratio ^C
H8 Samples				
1 7 3 8 and 9	0 36 56 70 and 77	70 74 72 72	16 18 8.2 14	0.23 0.24 0.11 0.20
E9 Samples				
4 5 6 10 11	0 8 56 74 94	83 83 71 73 78	11 4.3 8.2 7.2 9.7	0.14 0.05 0.12 0.10 0.12

a. Initial U concentration in the core was 65 wt%, in a control rod fuel assembly 64 wt%, and in a fuel rod 71 wt%.

b. Initial Zr concentration in the core was 18 wt%, in a control rod fuel assembly 17 wt%, and in a fuel rod 18 wt%.

c. The Zr/U ratios for a control rod fuel assembly and the core are 0.26 and 0.27, respectively. A fuel rod has a ratio of 0.25.

Zr concentrations (wt%) and Zr/U ratios for the ten recombined bulk samples. Relatively consistent U concentrations (±7%) and varying Zr/U ratios (1.e., varying Zr concentrations) are apparent. At the H8 location, the elemental concentrations and Zr/U ratios near the surface of the debris bed and at 36 cm into the debris bed are similar. Also, the Zr/U ratios are similar to those initially found in a control rod fuel assembly (1.e., 0.26). At 56 cm into the debris bed (Sample 3), the concentration of U is similar to the concentrations higher in the debris bed; however, the Zr/U ratio is approximately half of the ratios for the core and fuel rods, suggesting depletion (~58%) of Zr has occurred in this sample. The Zr depletion is probably caused by melting and relocation. The results are evaluated as they relate to the condition of the core in Section 4.3. The concentrations and ratios for the combined Samples 8 and 9 are similar to those obtained near the surface of the debris bed.

The Zr/U ratios at the £9 location range from 0.14 to 0.05, indicating depletion of Zr by 46 to 80%. There is significant depletion of Zr at all depths at the £9 location, whereas only one sample from H8 (Sample 3) showed significant Zr depletion. The data suggest that significant Zr depletion has occurred in the debris bed.

Tables 15 and 16 summarize from Appendix D the weight percent of U present in each particle and aliquot. Tables 17 and 18 list the Zr/U ratios for particles and aliquots obtained from the H8 and E9 samples, respectively. The results for the particles and aliquots show a much wider range of U concentrations (O to ~100% U) than do the recombined bulk samples (70 to 80% U), partly because the particles were chosen for specific characteristics (e.g., cladding, fuel pellet, etc.). Observations concerning these results and their significance are as follows:

few particles are completely U or Zr. The results indicate that little U is present without significant amounts of associated Zr. Little intact fuel material (88% U in UO₂ fuel or 71% U in a fuel rod) is present, and significant disruption of fuel rods is indicated.

TABLE 15. URANIUM CONTENT OF PARTICLES AND ALIQUOTS FROM THE H8 CORE LOCATION^a (wt%)

Particle Size Fraction (µm)	Particle/Aliquot	Sample 1	Sample 7	Sample 3	Sample 8	Sample 9
>4000	А	b	59,	38	72	40
>4000	В	83	b	b ⁰	81_	55
>4000	С	9.8	b	D	∿100g	78
>4000	D	26	b	48	b	11
>4000	E	17.8	b	72	√100 ^d	7 2
1680-4000	F	57	72	∿100d	55	67
1680-4000	G	57	52	29	12	13
1680-4000	Й	47	52	45	69	42
1000-1680	Ţ	0	b	56	22	14
1000-1680	j	64	40	11	63	45
1000-1680	K	57	68	48	81	b
707-1000	Aliquot	52	37	58	28	51
	Aliquot	53	35	42	42	b
297-707		45	b	37	32	67
149-297	Aliquot	41	52 _.	35	48	34
74-149	Aliquot	36	b	32		34
30-74	Aliquot	C	19	32 b	43 b	17
20-30	Aliquot Aliquot	c	13	- -	49	50
<20	ATIQUOL				73	30

a. In order of depth into the debris bed.

b. Not reported because of analysis problems.

c. Not analyzed.

d. The analysis results indicate the presence of other components. Also the uncertainty is large for these analyses indicating that this is oxidized uranium.

TABLE 16. URANIUM CONTENT OF PARTICLES AND ALIQUOTS FROM THE E9 CORE LOCATION® (wt%)

Particle Size Fraction (µm)	Particle/Aliquot	Sample 4	Sample 5	Sample 6	Sample 10	Sample 11
		04	62	~100 ^b	C	~100b
>4000	A	94	85	V100	88	C
>4000	В	58	57	Ŏ	71	~100b
>4000	C	51	5/ 00	8.1	61	~100b
>4000	D	c	90 55 c		65	56
>4000	E	88		0	05	30
1600 4000	•		C	0	63	82.
1680-4000	r		c	90	81	~100 ^b
1680-4000	G		c	71	68	~100b c
1680-4000	H			71	•	
1000 1600	7		17	11	35	71
1000-1680	i .		<u></u> c	13	54	63
1000-1680	 		c	70	~100 ^b	96
1000-1680	K			, 0	100	-
707-1000	Aliquot		31	55	64]4
	Aliquot			44	38	a
297-707				31	33	71
. 149-297	Aliquot			32	17	40
74-149	Aliquot			28	37	34
30-74	Aliquot				47	22
20-30	Aliquot					~100b
<20	Aliquot	~~				

a. In order of depth into the debris bed.

b. The analysis results indicate the presence of other components. Also, the uncertainty is large for these analyses, indicating that this is oxidized uranium.

c. Not reported because of analysis problems.

d. Not analyzed.

TABLE 17. ZR/U CONCENTRATION RATIOS OF PARTICLES AND ALIQUOTS FROM THE H8 CORE LOCATIONa

Particle Size Fraction						
<u>(µm)</u>	Particle/Aliquot	Sample 1	Sample 7	Sample 3	Sample 8	Sample 9
>4000	Α	b	2.2 E-1	6.0 E-2	1.1 E-1	1.5 E-1
>4000	В	1.5 E-2	b	C	9.8 E-2	4.4 E-1
>4000	С	1.5 E-1	þ	b	5.7 E-1	6.6 E-2
>4000	D	6.1 E-2	b b	2.0 E-2	b	6.7 E-1
>4000	E	3.7	n	3.2 E-2	5.1 E-1	8.7 E-1
1680-4000	. F	2.4 E-1	4.3 E-2	5.1 E-2 1.4 E-1	5.4 E-1 8.0 E-1	2.9 E-1 4.3 E-2
1680-4000	G	2.4 E-1	2.6 E-1		1.2 E-1	1.2 E-1
1680-4000	Н	7.8 E-1	7.4 E-3	2.2 E-1	1.2 = 1	1.2 6-1
1000-1680	I	c	b	1.5 E-1	2.2 E-1	2.0
1000-1680	J	2.3 E-1	6.2 E-2	2.4	7.9 E-1	1.2 E-1
1000-1680	K	2.3 E-1	2.1 E-1	1.9	2.5 E-1	
707-1000	Aliquot	2.3 E-1 1.9 E-1	2.3 E-1 7.2 E-1	3.4 E-1 1.9 E-1	1.1 9.3 E-2	1.18 b
297-707	Aliquot	3.2 E-1	7 • 2	4.7 E-2	5.8 E-1	5.4 E-1
149-297	Aliquot	7.3 E-1	8.4 E-1	9.2 E-1	5.4 E-1	4.4 E-1
74-149	Aliquot	7.9 E-1	0.7_b	1.1	6.4 E-1	5.8 E-1
30-74	Aliquot		d	'•' b	0.4 L- ,	2 g
<30	Aliquot	9.3 E-1 d	9.2 E-1	d	b	5.4 E-1
20-30	Aliquot	d	9.2 E-1	2.7 E-1	d	2.9 E-1
<20	Aliquot			Z./ E-1	** ** −	2.3 L-1

a. In order of depth into the debris bed.

b. Not reported because of analysis problems.

c. All zirconium or uranium; the other element was not detected.

d. No particle size analysis was performed for this size group.

TABLE 18. ZR/U CONCENTRATION RATIOS OF PARTICLES AND ALIQUOTS FROM THE E9 CORE LOCATION®

Particle Size Fraction (µm)	Particle/Aliquot	Sample 4	Sample 5	Sample 6	Sample 10	Sample 11
>4000 >4000 >4000 >4000 >4000	A B C D E	1.4 E-1 3.1 E-2 2.4 E-2 b 3.0 E-2	1.4 E-1 4.9 E-2 6.8 E-2 4.6 E-3	2.4 E-2 c b 1.0 E-2	7.8 E-2 4.6 E-3 4.4 E-3 1.2 E-1	8.5 E-2 b 4.5 E-3 1.8 E-2 5.4 E-3
1680-4000 1680-4000 1680-4000	F G H	a 	b b	c 5.5 E-3 5.8 E-3	6.3 E-1 4.6 E-3 5.7 E-2	1.1 E-1 3.0 E-3
1000-1680 1000-1680 1000-1680	I J K	 	4.9 E-1 b b	2.0 4.2 1.1 E-1	6.8 E-2 1.6 E-1 5.8 E-3	3.0 E-1 3.2 E-3 3.1 E-3
<1000 707-1000 297-707 149-297 74-149 30-74 <30 20-30	Aliquot	 	9.2 E-2 	2.8 E-1 3.9 E-1 2.1 E-1 1.0	3.3 E-1 1.7 E-1 6.8 E-1 7.4 E-1 4.4 E-1 d 3.3 E-1	4.2 b 1.8 9.7 E-1 1.4

a. In order of depth into the debris bed.

Not reported because of analysis problems.

c. All zirconium; no uranium was detected.

d. No particle size analysis was performed for smaller sizes.

- Higher concentrations of Zr are present in the smaller particle 0 size fractions, with a maximum Zr/U atom ratio of approximately 1:2.5 for the smallest (<30 µm) particle sizes. This observation is consistent with previously reported results characterizing debris from fuel damage tests in the Power Burst Facility reactor of INEL. 23 The higher concentrations of Zr are probably due to fragmentation of the zircaloy cladding remnants and increasing oxygen content (oxidation) for the smaller particle sizes. The Zr/U atom ratios range from 1:7 to 1:2.5. An increase in Zr/U ratio with diminishing particle size is apparent in samples to a depth of 56 cm (Sample 3) at the H8 location (see Table 17). No measurable correlation between the Zr content and particle size is evident below this depth. The Zr content in the E9 samples (see Table 18) shows a particle size dependency which extends through 94 cm into the debris bed (Sample 11). The only exception is Sample 10 (E9. 74 cm), where the Zr content is about the same for all particle size fractions.
- 3.3.2.2 <u>Control Rod Materials (Ag-In-Cd)</u>. The Ag-In-Cd control rod materials (2749 kg) account for approximately 2.2 wt% of the core weight. They are important because they are highly volatile and likely to vaporize and condense into aerosols, a principal mode for transport from the core to the reactor coolant system. The three constituents of the control rods, in their relative order of volatility (most volatile to least volatile), are Cd, In, and Ag.

Tables 12 and 13 summarize the elemental analysis results for Ag, In, and Cd in the recombined bulk samples. Of these elements, only Ag was measurable in the recombined bulk samples. The measured Ag concentrations are generally similar for both the H8 and E9 samples, ranging from 0.13 to 0.19 wt%. This suggests a relatively homogeneous distribution in the rubble bed. These concentrations are equivalent to only 7.4 to 10% of the expected Ag concentration of the core. The measured detection limit values for Cd and In suggest significant depletion of both elements.

The particle and aliquot data also indicate that Ag is relatively evenly distributed (0.1 to 0.2 wt%) at the sampled locations, with some localized areas of higher concentrations. The higher concentrations in the smaller particle sizes suggest Ag may be present primarily as relatively small nodules or localized accumulations.

Although Cd and In were not measurable in the recombined bulk samples, the detection limits were below expected core average concentrations suggesting significant depletion. Ag-In-Cd were measurable in many of the particles and aliquots (see Appendix D) because of a lower detection limit associated with the sample size analyzed. Where measurable, Cd and In concentrations are generally higher (0.3 to 0.8 wt%) than Ag. The original concentration of In in a control rod fuel assembly is approximately 0.9 wt%. The number of samples where In was not measurable may be due to the ICP detection limit for In, which is approximately a factor of four higher than the limit for Ag (up to 0.6 wt% for these samples).

Cd was measurable in a larger fraction of the total number of samples than In, but in fewer than was Ag. The samples with measurable amounts of Cd are principally particle size fractions less than 1000 µm. The smallest size fractions have the highest Cd concentrations (0.2 to 0.5 wt%). The concentration of Cd in a control rod fuel assembly is approximately 0.3 wt%. The data suggest Cd retention is similar to In.

No correlation between concentrations of the three control rod constituents is apparent for the particles and aliquots. The higher concentrations of In and Cd suggest that these elements are forming accumulations in the core. For the smaller ($\leq 707~\mu m$) particle size fractions, there is an increase in concentrations for control rod constituents with decreasing particle size. The relative surface areas of the individual particle size fractions were calculated (Appendix G) to determine if a correlation exists between elemental concentration and particle size. A comparison of the increases in concentration of the Ag-In-Cd with the increase in surface area was made for the different

particle size fractions. No linear correlation with particle size was found; however, higher concentrations of the Ag-In-Cd materials appear to be associated with the smaller particles.

3.3.2.3. <u>Burnable Poison Rod Materials</u>. The two burnable poisons present in the core are $B_4C-Al_2O_3$ and $Gd_2O_3-UO_2$. Elemental constituents measured during analysis of the debris were Al, Gd, and B. Al and Gd were among the least abundant elements in the core, with original inventories of 214 and 13.5 kg, respectively. However, B was quite abundant, with approximately 172 kg from the poison rods and deposited amounts from the borated reactor coolant.

The results presented in Tables 12 and 13 indicate that Gd was measurable in most of the recombined bulk samples at concentrations of 0.04 to 0.08 wt%, whereas Al was measurable only in the two surface samples (Samples 1 and 4) at concentrations of 3.37 and 1.1 wt%.

Observations concerning these results and their significance are as follows:

Gd is relatively evenly distributed in the debris bed at 0 approximately 0.056 wt%. Only four fuel assemblies in the core [at locations H4, H12, D8, and N8 (see Figure 2)] contained $Gd_2O_3-UO_2$ poison rods. The Gd concentrations at E9 are similar to H8, although the distances from E9 and H8 to the nearest Gd rod are approximately 30 and 87 cm, respectively. These data indicate Gd has been well distributed and evenly mixed with material present in the debris bed. The Gd concentration is 4 to 7 times higher than would be expected if the Gd was evenly distributed in the entire core (~0.011 wt%). The measured concentrations, if extrapolated to the estimated mass of the debris bed (~20% of the core mass) would account for almost 100% of the original Gd inventory. These data suggest, along with the Zr and Ag results, that the composition of the core below the debris bed may be significantly different.

- High concentrations of Al are present at the surface of the 0 debris bed. This indicates the presence of a mechanism causing Al to accumulate at the surface rather than become evenly distributed within the debris bed. The measured concentrations for Al at the surface are 7 to 20 times greater than if the Al were evenly distributed in the debris bed (~1.7 mg Al/g Several mechanisms for the uneven distribution of Al have been suggested. Possible mechanisms are the deposition of $\mathrm{Al}_2\mathrm{O}_3$ or $\mathrm{AlO}(\mathrm{OH})$ powder in the surface of the debris during natural recirculation or possibly vaporization/ deposition of the Al on the in-core surfaces. Volatile Al could be produced by the reduction of AL₂O₃ by liquified zirconium. The results of the particle and aliquot analysis indicate there is a relationship between particle size and Al concentration, with generally higher concentration in the smaller particle size fractions.
- 3.3.2.4 <u>Structural Materials</u>. The principal structural materials measurable in the core debris are fe, Mn, Mo, Cr, Ni, Nb, and Sn. Table 19 lists average concentrations of these elements, ranges, and comparisons with average concentrations in the debris bed. No significant accumulations of any element were found in the debris, with the possible exception of Nb, which was expected because it has a high melting point (>2600 K). The results indicate that there has been significant depletion (50 to 75%) of structural materials from the debris bed.
- 3.3.2.5 <u>Tellurium</u>. Due to radioactive decay, no fission product Te radionuclides were measurable in the core debris grab samples. Therefore, elemental Te content was measured in an attempt to determine the concentration of fission product Te in the debris bed at the time of the accident.

Tables 12 and 13 indicate that Te concentrations were less than the ICP detection limit for the recombined bulk samples; however, Te was measurable in some particles and aliquots, but was near the ICP detection

TABLE 19. STRUCTURAL MATERIAL CONCENTRATIONS AND RANGES OF THE TMI-2 CORE DEBRIS GRAB SAMPLES

Element	Measured Average Concentration (wt%)	Range	Initial Core Average Concentration ^a (wt%)	Fraction of Core Inventory ^b
H8 Samples				
Fe	9.6 E-1	5 E-1 to 1.1	2.8	0.34
Mn	3.3 E-2	1.9 E-2 to 4.7 E-2	0.07	0.47
Cr	2.9 E-1	1.2 E-1 to 3.9 E-1	0.91	0.32
Мо	c	c	0.03	c
Ni	3.8 E-1	1.7 E-1 to 6.5 E-1	0.84	0.45
Nb	1.7 E-1d	0 to 1.7 E-1	0.05	3.1
Sn	c	c	0.29	c
E9 Samples				
Fe	4.6 E-1	2.1 E-1 to 8.3 E-1	2.8	0.16
Mn	1.7 E-2	1.4 E-2 to 2.1 E-1	0.07	0.24
Cr	1.2 E-1	6.3 E-1 to 2.1 E-1	0.91	0.13
Мо	c	c	0.03	c
Ni	2.9 E-1d	2.4 to 39	0.84	0.35
Nb	1.8 E-1d	1.0 to 2.7	0.05	3.6
Sn	C	c	0.29	C

a. A total core mass of 1.25 x $10^5\ kg$ and an even distribution of the core inventory from Table 11 is assumed.

b. Fraction of core inventory is calculated by dividing the measured concentration by the average core concentration.

c. Not detected.

limit. Table 20 lists which particles and aliquots contain Te and summarizes the measured Te concentrations. The results indicate that most (75%) of the materials containing Te were located near the surface of the debris bed. The calculated Te inventory in the core at the time of the accident was approximately 4.45 kg, of which 800 g was estimated to be Te impurity in stainless steel and 3.65 kg was stable fission product Te. Assuming a total core mass of 1.25×10^5 kg, the average Te concentration was 3.6×10^{-3} wt%. The measured concentrations range from 25 to 277 times the average core concentration, suggesting significant relocation of Te to the surface of the debris bed. The source of the Te (i.e., either natural or fission product) is not known; however, no accumulations of other stainless steel constituents are present in the surface of the debris bed except A1.

3.4 Radiochemical Analysis

Radiochemical analyses were performed on the recombined bulk samples, particles and aliquots from all ten samples, and on the ferromagnetic component of Sample 6. Gamma spectroscopy, fissile material, and ⁹⁰Sr analyses were performed on the recombined bulk samples. These same analyses and ¹²⁹I and fertile material content analyses were performed on the particles and aliquots. Only gamma spectroscopy analysis was performed on the ferromagnetic component of Sample 6, because this portion had to be recombined with the bulk sample to allow measurement of the average radionuclide concentrations.

Radionuclides were chosen for analysis based on their importance to nuclear safety issues, because they would provide information on chemical behavior (volatility, leachability, etc.) and fission product behavior that will help define mechanisms by which radionuclides are released from the reactor core during severe fuel damage accidents. Evaluations of the radiochemical analysis results were performed to (a) determine the average radionuclide composition of the debris bed, (b) compare measured radionuclide concentrations with calculated radionuclide inventories (using the ORIGEN-2 code), (c) correlate fission product concentrations and

TABLE 20. TELLURIUM CONCENTRATIONS OF PARTICLES AND ALIQUOTS FROM THE TMI-2 CORE DEBRIS GRAB SAMPLES

Particles and Aliquots Containing Tellurium	Tellurium ^a Concentration (wt%)
H8 Samples	
Particle 1B	0.5
Particle 1E	0.3
Particle 1F	0.4
Particle 1H	0.3
74–149 μm aliquot (Sample l)	0.8
30-74 µm aliquot (Sample 3)	0.8
>20 µm aliquot (Sample 8)	1.0
E9 Samples	
Particle 4A	0.5
Particle 4C	0.3
Particle 4E	0.5
Particle 5C	0.3
Particle 5I	9.0 E-2
Particle 6J	0.1
Particle 11A	0.8

a. The calculated average tellurium concentration for the original core is 3.6 x 10^{-3} wt%.

material types (e.g., zircaloy and structural materials), (d) estimate relocation of core materials based on the enrichment of individual samples, and (e) study the Cs release and settling characteristics of the core debris materials.

3.4.1 Summary of Results

Tables 21 and 22 list the measured radionuclide concentrations for the recombined bulk samples from the H8 and E9 core locations, respectively. Results are reported in µCi/g sample. Weights of the recombined bulk samples also are listed. Radionuclide concentrations of the insoluble portion of each recombined bulk sample, as determined by gamma spectroscopy are listed in Appendix E. As discussed in Section 3.3, less than 3% of any sample was insoluble, therefore, the radionuclide concentration of the insoluble portion is relatively insignificant when compared with the dissolved portion. The total uncertainty associated with the radionuclide analysis of the recombined bulk samples is approximately 10%. Factors contributing to the total uncertainty include (a) the presence of insoluble material, (b) uncertainties associated with the analytical methods, and (c) uncertainties associated with sample weight and volume measurements.

Appendix E presents results of the radiochemical analyses of the particles and aliquots. The radionuclides measured are the same as those listed for the recombined bulk sample in Tables 21 and 22. Results of the 129 I and fissile/fertile material content analyses also are included in Appendix E. Uncertainties associated with analysis of the particles and aliquots are generally 10%. However, as noted in Appendix E, uncertainties for some particles and aliquots are as high as 30%, due to either mass attenuation corrections for radionuclides measured using low energy (<500 keV) gamma rays (155 Eu is 30%), small (<5 mg) sample weights, or sample losses during dissolution.

Table 23 lists the results of the gamma spectroscopy analysis performed to determine the radionuclide retention by the ferromagnetic constituent of Sample 6.

TABLE 21. AVERAGE RADIONUCLIDE CONCENTRATIONS OF THE RECOMBINED BULK SAMPLES FROM THE H8 CORE LOCATION (μ C1/g sample decay corrected to April 1, 1984)

	Sample 1	Sample 7	Sample 3	Samples 8 and 9
<u>Radionuclide</u>	$(23.56 \text{ g})^{a}$	$(44.62 \text{ g})^{a}$	(50.10 g) ^a	(94.6 g) ^a
60 _{Co} 90 _{Sr}	4.99 ± 0.12 E+1 5.21 ± 0.23 E+3	3.93 ± 0.11 E+1 5.49 ± 0.33 E+3	6.44 ± 0.11 5.99 ± 0.24 E+3	3.41 ± 0.06 E+1 5.52 ± 0.22 E+3
106 _{Ru} 125 _{Sb}	4.92 ± 0.13 E+2 1.19 ± 0.03 E+2	3.61 ± 0.11 E+2 6.52 ± 0.28 E+1	7.56 ± 0.17 E+2 1.11 ± 0.04 E+2	3.80 ± 0.08 E+2 7.27 ± 0.23 E+1
1291d 134 _{Cs}	5.6 E-4C 6.73 ± 0.14 F+1	6.47 ± 0.14 E+1	3.8 E-4 5.22 ± 0.12 E+1	5.53 ± 0.08 E+1
137 _{Cs}	1.35 ± 0.01 E+3	1.38 ± 0.01 E+3	1.25 ± 0.01 E+3	1.30 ± 0.01 E+3
144Ce 154Eu	3.04 ± 0.21 E+3 5.49 ± 0.21 E+1	3.02 ± 0.21 E+3 5.30 ± 0.21 E+1	2.95 ± 0.20 E+3 4.67 ± 0.19 E+1	2.78 ± 0.01 E+3 4.65 ± 0.10 E+1
155Eս 235yb	1.00 ± 0.30 E+2 1.64 ± 0.59 E+1	9.4 ± 3.1 E+1 1.78 ± 0.09 E+1	8.3 ± 2.6 E+1 1.94 ± 0.10 E+1	8.4 ± 1.8 E+1 1.84 ± 0.02 E+1

a. Sample weight dissolved.

b. Analyses for 235 U were performed by the methods discussed in Section 2 and are reported in mg/g sample.

c. Results calculated from analysis of the particles and aliquots.

d. Analyses were performed for 129 I, however, safety restrictions made it necessary to reduce the sample size, thereby reducing the number of measurable results below the quantity required to calculate bulk sample concentrations.

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TABLE 22. AVERAGE RADIONUCLIDE CONCENTRATIONS OF THE RECOMBINED BULK SAMPLES FROM THE E9 CORE LOCATION (μ C1/g sample decay corrected to April 1, 1984)

Radionuc ilde	Sample 4 (5.11 g) ^a	Sample 5 (16.64 g) ⁸	Sample 6 (45.97 g) ^a	Sample 10 (57.44 g) ^a	Sample 11 (42.54 g) ^a
60 _{Co}	5.68 ± 0.11 E+1	4.68 ± 0.14 E+1	3.72 ± 0.12 E+1	3.25 ± 0.11 E+1	5.18 ± 0.14 E+1
90 _{Sr} 106 _{Ru}	6.33 ± 0.21 E+3	6.73 ± 0.34 E+3	5.07 ± 0.27 E+3	5.99 ± 0.28 E+3	4.97 ± 0.25 E+3
106 _{Ru}	7.12 ± 0.11 E+2	1.01 ± 0.02 E+3	5.46 ± 0.19 E+2	6.54 ± 0.17 E+2	5.65 ± 0.16 E+2
125 _{Sb}	8.13 ± 0.19 E+1	1.45 ± 0.03 E+2	1.01 ± 0.04 E+2	9.72 1 0.36 [+]	1.94 ± 0,04 E+2
125Sb 1291d	2.4 E-4C	5.9 E-4C	4.2 E-4C	0	a
134 _{Cs}	2.43 ± 0.06 E+1	1.09 ± 0.02 E+2	8.56 ± 0.02 f+1	6.69 ± 0.14 E+1	7.57 ± 0.18 E+1
137 _{Cs}	4.77 ± 0.02 E+2	2.44 ± 0.07 E+3	2.11 ± 0.01 E+3	1.56 ± 0.06 E+3	1.91 ± 0.01 E+3
144Ce	3.64 ± 0.18 E+3	3.05 ± 0.02 E+3	2.15 ± 0.22 E+3	2.58 ± 0.22 E+3	2.59 ± 0.22 [+3
154 _{Eu}	5.47 ± 0.17 E+1	4.88 ± 0.26 E+1	2.90 ± 0.19 E+1	4.14 ± 0.21 E+1	3.23 ± 0.18 E+1
155È	1.14 ± 0.36 E+2	9.8 ± 3.2 €+1	7.8 ± 2.6 E+1	8.8 ± 2.9 E+1	8.0 ± 2.6 E+1
235บู้ชื่	2.11 ± 0.21 E+1	2.29 ± 0.10 E+1	1.94 ± 0.07 E+1	1.96 ± 0.09 E+1	2.11 ± 0.11 E+1

a. Sample weight dissolved.

b. Analyses for 2350 were performed by the methods discussed in Section 2 are reported in mg/g sample.

c. Results calculated from analysis of the particles and aliquots.

d. Analyses were performed for ¹²⁹I, however, safety restrictions made it necessary to reduce the sample size, thereby reducing the number of measurable results below the quantity required to calculate bulk sample concentrations.

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TABLE 23. RADIONUCLIDE CONCENTRATIONS OF THE FERROMAGNETIC COMPONENT OF SAMPLE 6 ($\mu\text{C1/g}$, decay corrected to April 1, 1984)

	Particle Size Fraction (um)			
Radionuc lide	707-1000	297-707	149-297	74-149
	(0.25 g) ^a	(0.19 g) ^a	(0.025 g) ^a	(0.024 g) ^a
54Mn	7.4 ± 3.0 E-1	4.0 ± 1.2	3.4 ± 1.2	5.2 ± 1.2
60Co	2.01 ± 0.02 E+2	7.31 ± 0.07 E+2	6.18 ± 0.07 E+2	5.48 ± 0.06 E+2
106Ru	3.22 ± 0.54 E+2	1.39 ± 0.17 E+3	7.47 ± 0.20 E+2	6.46 ± 0.17 E+2
125Sb	2.35 ± 0.03 E+2	3.34 ± 099 E+2	3.23 ± 0.10 E+2	3.46 ± 0.09 E+2
134Cs	1.86 ± 0.35	8.79 ± 0.25 E+1	5.26 ± 0.22 E+1	6.72 ± 0.27 E+1
137Cs	3.61 ± 0.06 E+1	1.58 ± 0.01 E+3	1.02 ± 0.01 E+3	1.24 ± 0.01 E+3
144Ce	1.24 ± 0.22 E+2	7.4 ± 1.1 E+2	4.7 ± 1.0 E+2	4.31 ± 0.86 E+2
154Eu	1.55 ± 0.53	1.35 ± 0.24 E+1	6.7 ± 2.0	7.4 ± 1.8
155Eu	8.7 ± 2.9	3.3 ± 1.1 E+1	2.0 ± 0.6 E+1	2.3 ± 0.7 E+1

a. Weight of the ferromagnetic component.

3.4.2 Radionuclide Composition of the Core Debris

The radionuclide composition of the core debris was determined based on the radionuclide concentrations of the recombined bulk samples (listed in Tables 21 and 22). Results are discussed in the order of expected fission product volatility.

fission products are categorized in Table 24 by volatility of the chemical group and element. This grouping is consistent with the WASH-1400 characterization. Some possible chemical compounds for each element are also summarized, and the boiling temperatures of elements and some compounds are listed. Radionuclides for which analysis of the core debris grab samples were performed also are indicated.

The high-volatility fission product groups (I, II, III, and IV.a) shown in Table 24 contain the noble gases, halogens, alkali metals, and heavy chalcogens, and are characterized by boiling points less than 1600 K for the elemental forms as well as for the listed oxide compounds.

The medium-volatility fission products are characterized by boiling points less than 3100 K (UO₂ melting). These materials include fission product elements from the Group VA metals, alkaline earths, some of the rare earths, and actinides. The volatility of elements in this classification is strongly dependent on chemical form. For example, BaO has a low volatility, BaH₂ has a medium volatility, and BaO₂ has a high volatility. The chemical form of Ba will depend primarily on fuel chemistry, the oxidizing potential in the reactor coolant system, and temperature.

The low-volatility materials include fission product elements from the noble metals, the remaining rare earths and actinides, tetravalents, and early transition elements. Generally the oxides of these elements also have low volatilities; however, LaO and some of the noble metal oxides are significantly more volatile than the element.

TABLE 24. CORE MATERIAL VOLATILITY GROUPS

WASH-1400 Group Number	Chemical Group	Element	Boiling ^a Temperature (K)	Volatility	Possible Compounds	Boiling ^a Temperature (K)	Volatility	Analysis Performed
Fission Pr	oducts							
I	Noble gases	Kr ^b Xe ^b	121 166	High High				
II	Halogens	Br I	332 458	High High	CsBr ^b CsI ^b HI	1573 1553 238	High High High	X
111	Alkali metals	Rbb	973	High	Rb I b Rb 20 Rb 20	1577 1273	High High	X
		Csb	963	Ніgh	Rb202 Cs1b Cs0H Cs20 Cs202 Cs2U04 b	1553 1553 1350° 923	High High High High High	
IV.a	Heavy chalcogens	Se ^b	958	High	Se0 ₃	453	High	X
		⊺ eb	1215	Hîgh	Se02 Te02 Te202 silver-telluride iron-telluride zirconium telluride tin-telluride nickel-telluride chrome-telluride	613 ~1450 	High High 	
Fission Pro	ducts							
IV.b	Group VA metals	Spp	2023	Medium	Sb ₂ 0 ₃ ,	1823	Medium	X
V	Alkaline earths	Sr Ba	1657 1913	Medium Medium	SrO ^b BaH ₂ BaO ^b BaO ₂	1673 1673 1073	Low Medium Low High	X
VI	Rare earths	Eu Sm Pm	1870 2051 2733	Medium Medium Medium	Eu ₂ 0 ₃ b Sm ₂ 0 ₃ b Rm ₂ 0 ₃ b			
	Actinides	Am	2880	Medium	AmO ₂ b			

WASH-1400 Group Number	Chemical Group	Element	Boiling ^a Temperature (K)	Volatility	Possible Compounds	Boiling ^a Temperature (K)	Volatility	Analysis Performed
AI	Noble metals	Pdb Rhb	3413	Low	PdO	•• ;	**	X
		RhD	4000	Low	Rh0 ₂ Rh ₂ 0 ₃		,	
		Rub	4173	Low	Ru02 Ru04	**		
		Mob	4885	Low	M002 D M0203	√1300°	High	
		Tcb	5 150	Low	Mo03		••	
VII	Rare earths	Y	3611	Low	Y203 b	••	••	
		La	3730	Low	LaO	4473	Medium-High Low	x
		Ce	3530	Low	La ₂ 0 ₃ b Ce0 ₂	 		
		Pr	3485	Low	Ce ₂ O ₃ b PrO ₂ Pr ₂ O ₃ b	••		
		Nd	3400	Low	Nd203	••		
	Actinides	Np	4175	Low	NpO2 b		**	
•		P u Cm	3505 	Low Low	PuO2 b CmO2 b	~3570 	Fow	
	Tetravalents	Zrđ	4650	Low	ZrO ₂ b	5273	Low	
	Early transition	Nb	5015	Low	Nb02 b Nb205		Low	

a. Boiling temperature at 1 atm., data primarily from CRC Handbook of Chemistry and Physics, 56th Edition.

b. Probable chemical form of the fission product within the fuel.

c. NAA-SR-132.

d. Zirconium is both a fission product and a structural material.

e. Probable form of oxide released because of MoO_2 decomposition.

3.4.2.1 Cesium and Iodine.

129 I and 134,137 Cs are the most volatile radionuclides measurable in the core debris grab samples and have been released to the reactor coolant system in significant quantities. The 129 I concentrations listed in Tables 21 and 22 were obtained by weighting the results of the particle and aliquot analyses.

129 I analysis was performed on recombined bulk samples (Samples 1, 3, 4, 5, and 6). Subsequent safety restrictions made it necessary to reduce the sample sizes for Samples 7 through 11, thereby reducing the number of measurable results below the number required to calculate average bulk sample concentrations. The 129 I results for the five measured samples are mutually consistent, suggesting relatively homogeneous 129 I concentrations in the debris bed. The estimated uncertainty for the calculated concentrations is at least 100%. Retention of 129 I in the fuel matrix and on structural materials is further evaluated in Section 3.4.3 and 3.4.4.

At the H8 sample locations, the concentrations of 137 Cs at all depths are relatively consistent (within 5% of the average concentration of 1.32 E+3 μ Ci/g sample). At the E9 sample locations, however, the 137 Cs concentrations vary more widely. The 137 Cs concentration for Sample 4 (E9, surface) is a factor of 5 lower than any other 137 Cs concentration within the debris bed. These data indicate a significant change in concentration at the surface of the debris bed. The concentration is highest for Sample 5 (E9, 8 cm) and generally decreases with increasing depth, suggesting the presence of a concentration gradient.

Comparing the measured ¹³⁷Cs concentrations at the H8 and E9 locations suggests significant differences in fission product behavior at the two locations:

The ¹³⁷Cs concentrations are higher (from 18 to 85%) at E9 than at H8, suggesting radial differences in the radionuclide composition of the core. The data suggest that the H8 and E9 locations experienced different conditions during or after the accident.

- o Although the ¹³⁷Cs concentrations at H8 are consistent, a gradient is indicated at the £9 location. A 22 to 36% decrease in ¹³⁷Cs concentration from Sample 5 (8 cm) to Samples 10 and 11 (74 to 94 cm in the debris bed) suggests that areas deeper in the core may have different radionuclide concentrations.
- 3.4.2.2 Strontium and Antimony. 90 Sr and 125 Sb have relatively moderate volatilities as elements. However, as compounds, Sb_20_3 has a moderate volatility, but Sr0 has a low volatility. The concentrations of 90 Sr for all H8 samples are within 6% of the average concentration (5.55 E+3 μ C1/g sample). These data indicate that the distribution of 90 Sr in the debris bed at the H8 location is relatively consistent. Also, comparison of the fissile material content with 90 Sr concentrations suggests that 90 Sr retention in the fuel is rather uniform. Other examinations (not part of this report) have determined that small amounts of 90 Sr have leached into the reactor coolant. Retention of 90 Sr in the fuel is discussed in Sections 3.4.3 and 3.4.4.

At the E9 location, the 90 Sr concentrations are more variable (within 15% of the average concentration of 5.82 E+3 μ C1/g sample). However, the data indicate that 90 Sr is relatively evenly deposited in the debris bed.

At the H8 and E9 locations, the ¹²⁵Sb concentrations vary by about a factor of 2. The average concentration at E9 is approximately 23% higher than at H8, suggesting a radial gradient or variation in burnup between the two locations. No correlation with fissile material content is apparent for ¹²⁵Sb.

3.4.2.3 Ruthenium. Ru fission products are expected to be relatively nonvolatile as the metal but more volatile in the oxide forms (especially RuO_4). The ^{106}Ru concentrations listed in Tables 21 and 22 vary significantly (up to a factor of 2.8) at the H8 and E9 locations. The concentrations at E9 are generally higher than at H8. At H8, the concentrations range from 3.61 E+2 to 7.56 E+2 μ C1/g, whereas at E9 the

concentrations range from 5.46 E+2 to 1.01 E+3 μ Ci/g, indicating a radial concentration gradient. No correlation between concentration and depth is apparent. Comparison with fissile material (235 U) content indicates significant relocation of 106 Ru from the fuel. The 106 Ru/ 235 U ratios range from 20.6 to 39 at H8 and from 27 to 44 at E9, suggesting variations in 106 Ru retention in the fuel. Variations in burnup can affect this correlation; however, further evidence of 106 Ru relocation and evidence of a correlation between 106 Ru and structural material content is discussed in Section 3.4.4.

3.4.2.4 <u>Cerium and Europium</u>. 144 Ce and 154,155 Eu are refractory materials in the lanthanide group and were expected to have been retained significantly in the fuel. 144 Ce is discussed principally because comparison of the 144 Ce/ 154,155 Eu ratios indicates that the cerium and Eu behave similarly, with the exceptions of Samples 6 and 11 which have higher than expected ratios.

The ¹⁴⁴Ce concentrations at H8 are similar (within 9%) at all depths. Correlations with fissile material content indicates higher ratios (12 to 22%) near the surface of the debris bed.

There is an apparent gradient (~29%) in the 144 Ce concentrations at the E9 locations, from high values at the surface to lower values at 94 cm into the debris bed. Correlation with fissile material content indicates higher 144 Ce/ 235 U ratios (~28%) near the surface of the debris bed. An explanation for the variation in ratios is discussed in Section 4.3.

3.4.2.5 <u>Cobalt</u>. 60 Co is an activation product of Co present in stainless steel and Inconel. The data suggest variations in concentrations by factors of 1.8 and 7.7 at H8 and E9, respectively. 60 Co concentrations can provide some indication of the structural material content of the core debris grab samples.

3.4.3 Comparison with ORIGEN-2 Predicted Radionuclide Concentrations

The measured radionuclide concentrations of the recombined bulk samples were compared with concentrations predicted by the ORIGEN-2 computer code in order to assess retention of radionuclides in the core and fuel matrix.

Two methods were used to estimate normalized radionuclide retention in the core materials:

- The measured radionuclide concentrations in the core debris grab samples were extrapolated to the weight of all materials present in the core (1.25 \times 10^5 kg) and ratioed to the whole core fission product inventories. This analysis assumes the debris bed is representative of the core as a whole.
- The radionuclide concentrations were also normalized to the amount of U present in each bulk sample (see Tables 12 and 13) and ratioed to the predicted average fission product concentration per gram of U. This analysis assumes that the U contains the radionuclide inventory and that the sample has an average burnup.

Appendix f presents the average radionuclide concentrations per gram of U, as predicted by a nodular ORIGEN-2 analysis. Average predicted radionuclide concentrations are used because the quantities of fission products associated with each gram of U varies with 235U enrichment and burnup (total neutron exposure). Figures F-1 through F-7 in Appendix F show the variations in burnup and enrichment before the accident as functions of radial position and depth in the core. Burnup of the reactor fuel and, consequently, fission product concentrations predicted by ORIGEN-2 vary from the average concentration by a factor of approximately 2 for the TMI-2 core. Therefore, comparing normalized measured to predicted concentrations is valuable principally for intercomparison purposes. However, a measure of the accuracy of the comparison is obtained by

comparing retention of the refractory materials (e.g., ¹⁴⁴Ce) with predicted concentrations to obtain an effective measure of retention. The ¹⁴⁴Ce results indicate retention greater than 100%, suggesting that the samples were obtained from locations with higher than average burnup. The uncertainty associated with each analysis method is addressed below.

Table 25 summarizes the normalized retention of radionuclides in the core, based on extrapolating the recombined bulk sample concentrations to the total core weight. With few exceptions, retention follows the order of volatility of the fission products discussed in Section 3.4.2. The $^{129}\mathrm{I}$, $^{134}\mathrm{Cs}$, and $^{137}\mathrm{Cs}$ radionuclide concentrations indicate the lowest retention, ranging from 20 to 30% of the core inventory at both the H8 and E9 locations. These data suggest significant (70 to 80%) release of these fission products from the debris bed. Measurements of $^{137}\mathrm{Cs}$ released to other portions of the reactor account for approximately 40% of the core inventory of $^{137}\mathrm{Cs}$.

Significant variations in retention of ¹⁰⁶Ru are indicated at the H8 and E9 locations. Although the ranges overlap for the two locations, average retention varies by approximately 25%, suggesting significantly more depletion at the H8 sample locations.

The measured retentions of 90 Sr and 125 Sb in the debris bed differ significantly, although both elements have similar volatilities (the expected compounds do not). The results indicate 90 Sr is significantly retained in the debris bed at both locations, as predicted. The calculated retentions are greater than 100% and are probably due to higher than average burnup at the sampled locations. For 125 Sb, however, the average measured retention is significantly less (21 to 31%) at both locations, indicating behavior similar to 137 Cs. No explanation is apparent for the low concentrations of 125 Sb; however, Sb appears to be scavenged in some cases by structural material (see Section 3.4.4.2) and may have been transported to other locations in the reactor core (see Section 4).

TABLE 25. RADIOMUCLIDE RETENTION IN THE THI-2 CORES

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*	•	H8 Samples			E9 Samples			
Rad lonuc 11de	Total Core Inventory ^b (C1)	Average <u>Concentration^C</u>	Fraction of Inventory Retained ^d (%)	Range*	Average <u>Concentration</u> C	Fraction of Core Inventory Retained ^d (%)	Range*	
90 _{Sr}	6.62 E+5	5.55 E+3	105	98-113	5.82 E+3	110	94-127	
106Ru	1.15 E+5	4.97 E+2	54	39-82	6.97 E+2	76	59-110	
125SP	3.70 E+4	9.20 E+1	31	22-40	6.23 E+1	21	4.9-34	
129]	2.29 E-1	4.70 E-4	26	21-31	4.17 E-4	23	13-32	
134Cs	3.70 E+4	5.99 E+1	20	18-23	1.23 E-1	24	8.2-37	
137Cs	7.60 E+5	1.32 E+3	55	21-23	1.70 E+3	28	7.8-40	
144Ce	2.75 E+5	2.95 E+3	134	126-138	2.80 [+3	127	98-165	
154Euf	6.38 E+3	6.03 E+1	98	91-108	4.12 [+]	81	57-107	
155 _{Eu} f	1.61 E+4	9.02 E+1	70	64-78	9.16 E+1	71	60-88	

a. Radionuclide retention is calculated assuming the core is homogeneous and the chemical composition is similar throughout the core.

- b. The total core fission product inventories (in Ci), as calculated by ORIGEN-2, are listed in Appendix F.
- c. Average concentration is listed in pC1/g debris; decay is corrected to April 1, 1984.
- d. Fraction of core inventory retained is calculated by extrapolating the average radionuclide concentration to the total core mass.
- e. The range is for the miminum and maximum calculated fractions of core inventory retained for all H8 or E9 samples.
- f. The predicted concentrations for \$154Eu and \$155Eu have relatively large associated uncertainties (20-30%) because of uncertainties in the production of these radionuclides as extrapolated to the entire core.

The refractory radionuclides (144Ce, 154Eu, and 155Eu) were significantly retained in the core debris at both the H8 and E9 locations. The 144Ce concentrations are significantly greater than 100% (i.e., ~130%). The probable explanation is that the core debris grab samples were obtained from locations with higher than average burnup. The uncertainty associated with radionuclide retentions as determined from an evaluation of the 144Ce results would suggest a maximum uncertainty of ~67% for the sample with the highest concentrations above the core average to a mean uncertainty of 34%. These data would cause a reduction in the calculated retention by 34% or 67%, respectively. This calculation uses a gross 15% uncertainty for the measurement results. However, this technique is considered less accurate than other methods, as the core must be assumed to be homogeneous and chemically similar.

To better evaluate retention in the fuel and, as previously discussed, provide another estimate of radionuclide retention in the core debris, the measured radionuclide concentrations were normalized to the measured U content for each sample (see Section 3.3) and compared to the average fission product concentrations calculated by ORIGEN-2. Table 26 summarizes the normalized retention of radionuclides in the core debris, based on fission product concentrations normalized to U content. Also included is the effective enrichment determined for each sample, based on the fissile material content from Tables 21 and 22 and U content from Tables 12 and 13. The enrichment results indicate significant variations from the original U enrichments at H8 (2.64%) and E9 (1.98%). At H8, some of the measured enrichments (i.e., Samples 7, 8, and 9) are similar to the original enrichments. However, at E9 the enrichments are significantly higher than the original 1.96% enrichment, indicating significant fuel mixing. Original burnup at E9 ranged from 1931 to 4485 MWd/MTU, whereas at HB the burnup ranged from 2057 to 6213 MWd/MTU. Although H8 had a higher original burnup, the data indicate significant relocation. The fractions of the core inventory retained are similar to those listed in Table 25. The uncertainty associated with this analysis ad determined from an evaluation of the Ce results would indicate a maximum uncertainty of

TABLE 26. RADIONUCLIDE RETENTION NORMALIZED TO URANIUM CONTENTA (% of core inventory)

	Calculated Core Average		H8 Samples ^C			[9 Samples ^C				
Radionuc lide	Concentration (C1/q U)	1	_3_	_3_	8 and 9	4_	_5_	6	10	_11_
905r 106Ru 125Sb 1291 134Cs	8.12 E-3 1.41 E-3 4.53 E-4 2.81 E-9 4.53 E-4	91 50 37 28 21	91 35 19 d 19	102 74 34 19 16	94 37 22 d	94 61 22 10 6.4	100 86 3.8 25	88 55 32 21 27	101 63 29 d 20	79 52 1.8 d
137 _{Cs} 144Ce 154Eu 155Eu 235Ue/238U (%)	9.32 E-3 3.37 E-3 7.82 E-5 1.98 E-4	21 128 100 72 2.3	20 121 92 64 2.4	18 121 82 58 2.7	19 114 82 59 2.5	6.2 130 84 69 2.5	32 110 75 60 2.8	32 90 52 56 2.7	23 104 72 61 2.7	26 99 53 52 2.7

a. Chemical analyses indicate there are no significant concentrations of structural components and, consequently, the calculated retention is representative of the debris bed without significant structural contamination.

- b. Calculated concentrations are from Appendix F and are decay corrected to Arpil 1, 1984.
- c. Uranium content used for normalization from elemental content in Section 3.
- d. Not reported.
- e. 235U values are from Table E-11, and total uranium from Tables 12 and 13.

34% with a mean value of 20%. Therefore, the measured retentions could conceivably be reduced by these fractional percentages (e.g., average 137 Cs retention would be reduced from 22% to 14.4%).

Table 27 compares the fractions of core inventory retained, calculated by extrapolating the measured fission product concentrations to the weight of all core components and normalizing to the quantity of fission products present in a gram of U. Retention calculated by either method is similar; however, the calculated retention based on the quantity of fission products in a gram of U is lower (10 to 15%). This method probably is more accurate, because the quantity of structural material in the debris bed to which extrapolations using core weight are made is not well known.

3.4.4 Fission Product Association With Zr and Structural Materials

Fission product association with Zr and structural materials indicates release of fission products from the fuel and subsequent association (by vaporization/deposition, chemical reaction, etc.) with other materials in the core region. Data used in this evaluation are from the gamma spectroscopy analysis of the ferromagnetic components of Sample 6 and from analysis of particles and aliquots that are predominantly U, Zr, or structural material.

3.4.4.1 Ferromagnetic Material Analysis. Table 28 lists the gamma spectroscopy analysis results from the measurable ferromagnetic components of Sample 6. These data provide an indication of the fission product concentrations associated with structural materials. Table 28 compares radionuclide concentrations of the ferromagnetic components with those of the recombined bulk samples. The data can be divided into two distinct groups of radionuclides. The first group, including 60 Co, 106 Ru, and 125 Sb, generally show concentrations higher than those of the average recombined bulk samples, whereas the concentrations of the second group, 134 Cs, 137 Cs, 154 Eu, and 155 Eu, are lower than the average concentrations. The 60 Mn (not measurable in the recombined bulk samples) and 60 Co are neutron activation products of stainless steel and are

TABLE 27. COMPARISON OF RADIONUCLIDE RETENTION CALCULATED BY DIFFERENT METHODS
. (% of core inventory)

	H8 San	ples	E9 Samples		
Radionuc 11d	Extrapolation e to Core Weight ^a	Extrapolation to a Gram of Uranium ^b	Extrapolation to Core Weight ^a	Extrapolation to a Gram <u>of Uranium^b</u>	
90sr 106Ru 125Sb	105	95	110	92 63	
luoka	54	49	76	63	
125 _{Sb}	31	28	21	18	
1291	26	23	23	19	
134Cs	20	18	24	19 21	
137 _{Cs}	22	20	28	24	
144Ce	134	121	127	107	
154Eu	98	89	81	67	
155 _{Eu}	70	63	71	60	

a. Results taken from Table 25.

b. Average concentrations obtained from Table 26.

TABLE 28. COMPARISON OF THE FERROMAGNETIC/AVERAGE RADIONUCLIDE CONCENTRATIONS OF SAMPLE 6ª

	Particle Size Fraction (µm)						
Radionuclide	707-1000 Aliquot	297-707 <u>Aliquot</u>	149-297 <u>Aliquot</u>	74-149 <u>Aliquot</u>			
54 _{Mn}	a	a	a	a			
60°C0	5.4	19.6	16.6	14.7			
106 Ru	0.59	2.6	1.4	1.2			
125 Sb	2.3	3.3	3.2	3.4			
134 _{Cs}	0.02	1.0	0.6	0.79			
137 _{Cs}	0.02	0.75	0.48	0.59			
144 Ce	0.06	0.34	0.22	0.20			
154Eu	0.06	0.47	0.23	0.26			
155 Eu	0.30	1.1	0.69	0.79			

a. This table lists the ratios of the radionuclide concentrations measured in the ferromagnetic component of Sample 6 to the recombined bulk sample average concentration. High ratios indicate accumulations of the radionuclide in the ferromagnetic component.

expected to be measurable in the ferromagnetic material fraction. The 106 Ru and 125 Sb, however, are fission products, and the data suggest that these radionuclides tend to be released from the fuel and become associated with structural materials. These data are further evaluated in the discussion in Section 3.4.4.2.

The average or lower than average concentrations of \$^{134}\text{Cs}\$, \$^{137}\text{Cs}\$, \$^{144}\text{Ce}\$, \$^{154}\text{Eu}\$, and \$^{155}\text{Eu}\$ were expected. The \$^{134}\text{Cs}\$ and \$^{137}\text{Cs}\$ are lower than expected because they may have been significantly leached into the coolant since the accident (see Reference 28) or were not strongly attracted to the structural materials. The low concentrations of \$^{144}\text{Ce}\$, \$^{154}\text{Eu}\$, and \$^{155}\text{Eu}\$ were expected because these materials tend to remain in the fuel.}

3.4.4.2 <u>Particle and Aliquot Analysis</u>. The analysis results of the particles and aliquots provide further indications of the retention of fission products by U, Zr, and structural materials. Chemical and radiochemical analyses were performed on each particle and aliquot studied. Table 29 lists the fission product concentrations for those particles that were principally U or Zr.

As shown in Table 29, ^{129}I and $^{134,137}Cs$ appear to be retained in different amounts between the U and Zr particles. ^{137}Cs is associated more with U than with Zr, whereas ^{129}I is associated to a greater extent with the Zr particles.

Table 30 summarizes the correlations of the particle and aliquot results shown in Figures 32 through 41. No correlations were observed between \$137_{CS}\$ or \$129_{I}\$ and uranium for the particles and aliquots (Figures 32 and 33). There is no apparent correlation between the \$137_{CS}\$ and \$129_{I}\$ concentrations [i.e., no similarity in the ratios of these radionuclides (Figure 34 and Table 29)]; however, the data in Table 29 are for a limited number of particles with concentrations ranging by a factor of 100, which may bias the average concentrations.

TABLE 29. COMPARISON OF FISSION PRODUCT CONCENTRATIONS ASSOCIATED WITH PARTICLES WHICH WERE PRINCIPALLY URANIUM OR ZIRCONIUM

	Principa	lly Uranium Bearing <u>Particles^a</u>	Principally Zi	Zr Rich	
Radionuclide	Average (µC1/g)	Range (µC1/g)	Average (μC1/g)	Range (μC1/g)	(μC1/g) U Rich (μC1/g)
60Co	1.2 E+1	0 - 70.4	1.45 E+1	1.2 - 5.9 E+1 2.6 E+1 - 1.65 E+2 1.3 - 1.16 E+2 6.9 - 3.05 E+2 2.2 E-5 - 4.8 E-3	1.2
90Sr	4.95 E+3	1.4 E+1 - 1.23 E+4	1.21 E+2		2.4 E-4
106Ru	1.01 E+3	5.40 E+2 - 1.73 E+3	3.3 E+1		3.3 E-2
125Sb	8.55 E+1	3.86 E+1 - 1.49 E+2	1.01 E+2		1.18
1291	6.38 E-4	3.5 E-5 - 1.6 E-3	1.20 E-3		1.88
134 _{Cs}	6.12 E+1	2.9 - 1.69 E+2	1.9 E+1	0.5 - 6.29 E+1	0.31
137 _{Cs}	1.82 E+3	5.3 E+1 - 4.38 E+3	5.40 E+2	1.17 E+1 - 1.24 E+3	0.30
144Ce	2.93 E+3	1.81 E+3 - 4.67 E+3	5.20 E+2 ^c	1.9 - 2.84 E+3	0.18
154 _{Eu}	3.79 E+1	1.58 E+1 - 7.87 E+1	7.7 E-1 ^c	0 - 0.98	2.0 E-2
155 _{Eu}	1.11 E+2	0 - 2.07 E+2	5.9 ^c	0 - 2.21 E+1	5.3 E-2

a. Particles were principally uranium with a few other contaminants. Particles 1B, 3F, 5D, 6A, 6G, 6H, and 7H were analyzed.

b. Particles were principally zirconium with little contaminants from other components. Particles 11, 38, 68, 6C, 6E, and 6F were analyzed.

c. Results were biased by Particle 6B.

TABLE 30. SUMMARY OF CORRELATIONS BETWEEN SELECTED RADIONUCLIDES, ELEMENTS, AND PARTICLE SURFACE AREA

<u>Figure</u>	Correlated Materials or Result	Comment	
32	137 _{Cs/U} content	No apparent correlation was observed between 137Cs and uranium content for particles or aliquots.	
33	129I/U content	No apparent correlation was found between 1291 and uranium content for particles or aliquots.	
34	129I/normalized particles surface area	There is a correlation between higher concentrations and smaller particle size, perhaps due to increased surface area. These results suggest vaporization/condensation on surfaces or possible reaction with nonfuel materials which have higher concentrations in the smaller particle size groups.	
35	137 _{Cs/} 129 _I	No apparent correlation between 137Cs and 129I was observed.	
36	106 _{Ru/N1} content	No correlation was observed for most particles and aliquots.	
37	125Sb/N1 content	No apparent correlation was observed.	
38	106 _{Ru/U} content	No obvious correlation is apparent.	
39	125 _{Sb/U} content	No correlation is apparent.	
40	125 _{Sb/} 129 _I	No correlation is apparent.	
41	144Ce/U content	No correlation is apparent.	

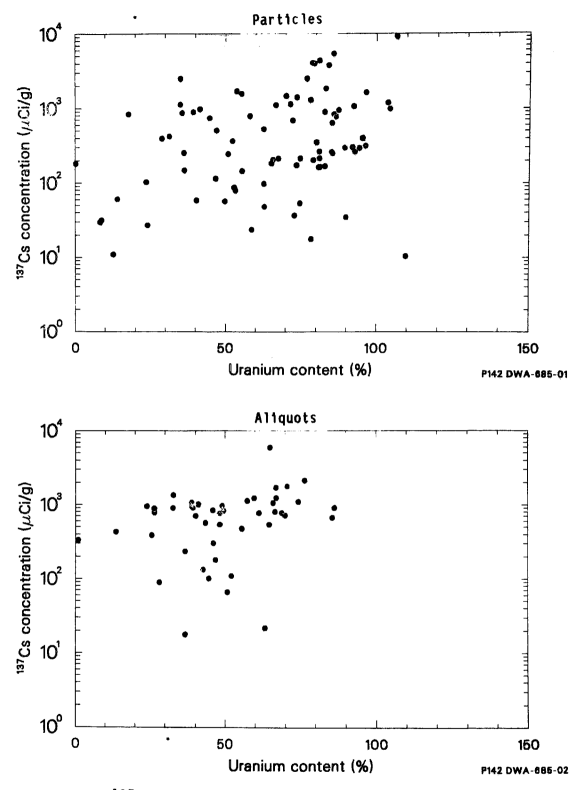


Figure 32. 137Cs/U content correlation for all particles and aliquots analyzed.

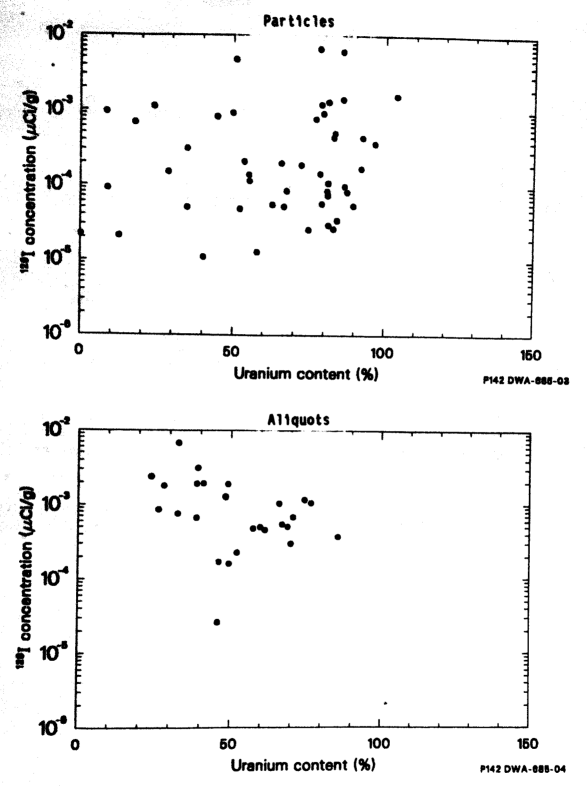


Figure 33. 129I/U content correlation for all particles and aliquots analyzed.

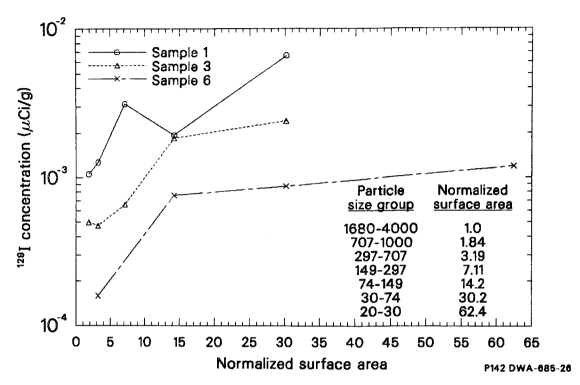


Figure 34. Relationship between ^{129}I concentration and surface area for Samples 1, 3, and 6.

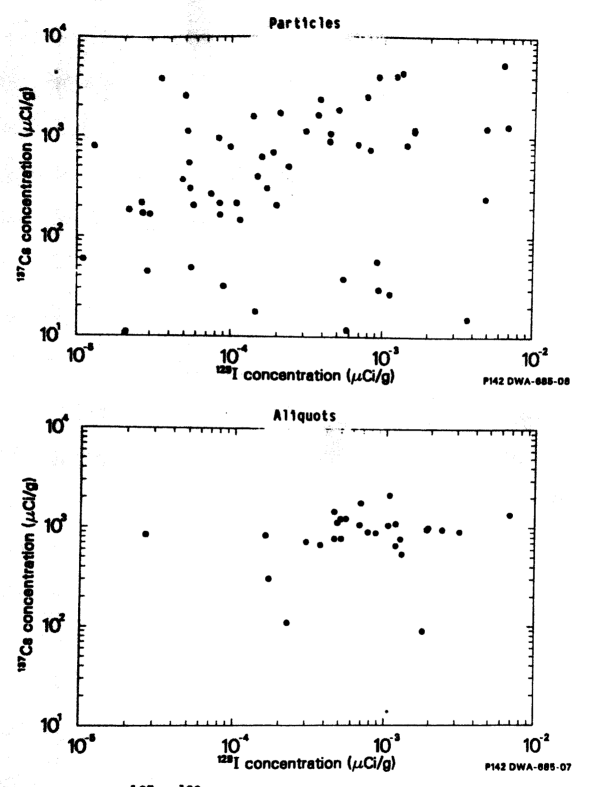


Figure 35. 137Cs/129I content correlation for all particles and aliquots analyzed.

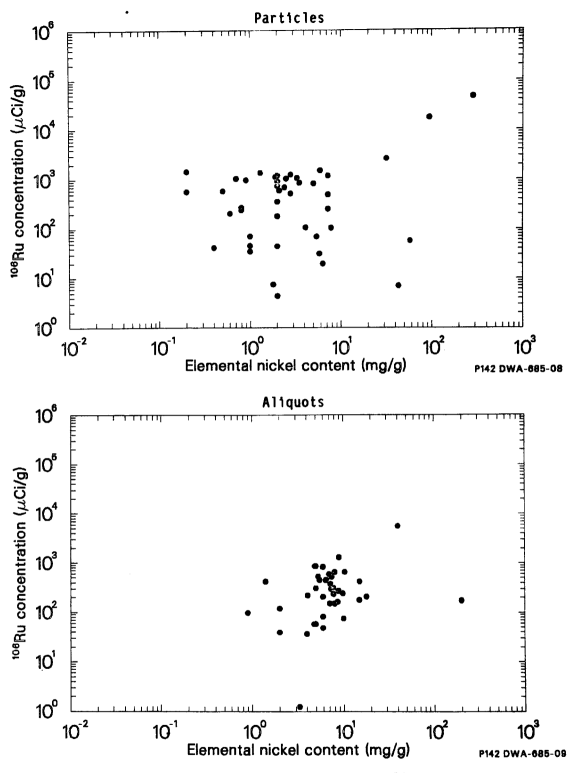


Figure 36. Correlation between Ni content and 106Ru for all particles and aliquots analyzed.

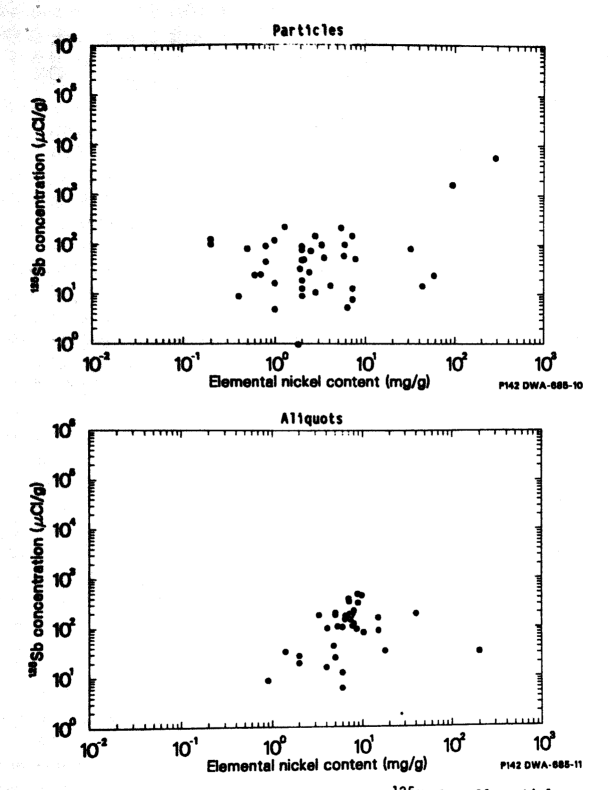


Figure 37. Correlation between Ni content and 125Sb for all particles and aliquots analyzed.

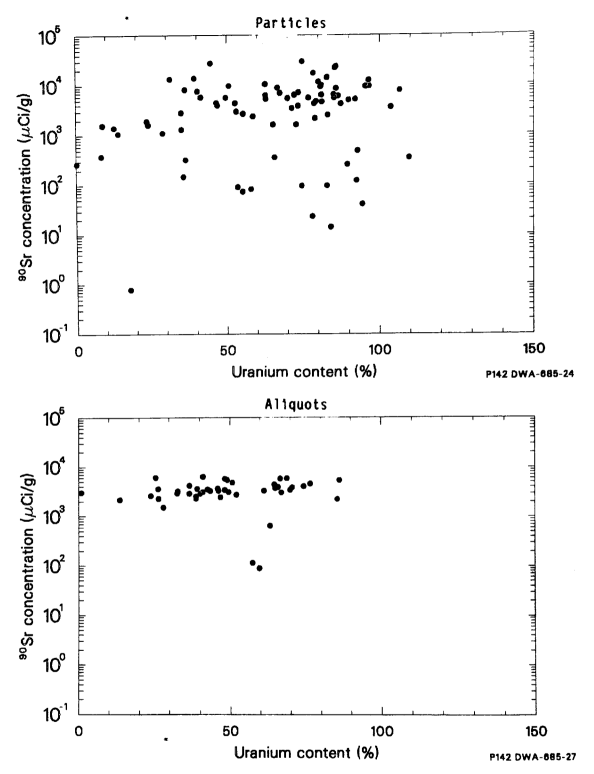


Figure 38. $^{90}\text{Sr/U}$ content correlation for all particles and aliquots analyzed.

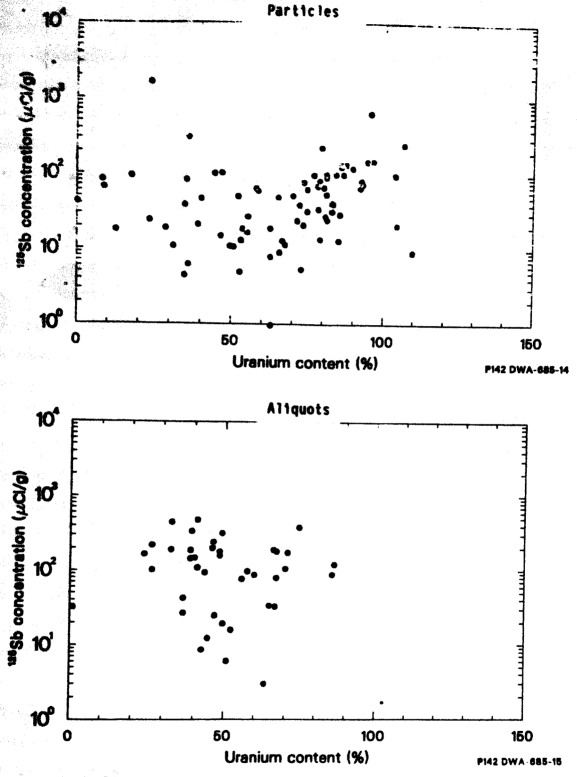


Figure 39. 125Sb/U content correlation for all particles and aliquots analyzed.

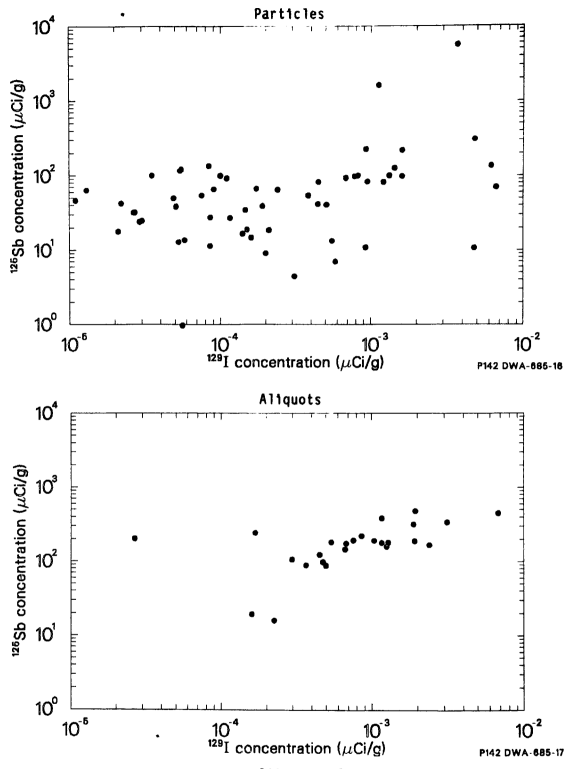


Figure 40. Correlation between $^{125}\mathrm{Sb}$ and $^{129}\mathrm{I}$ for all particles and aliquots analyzed.

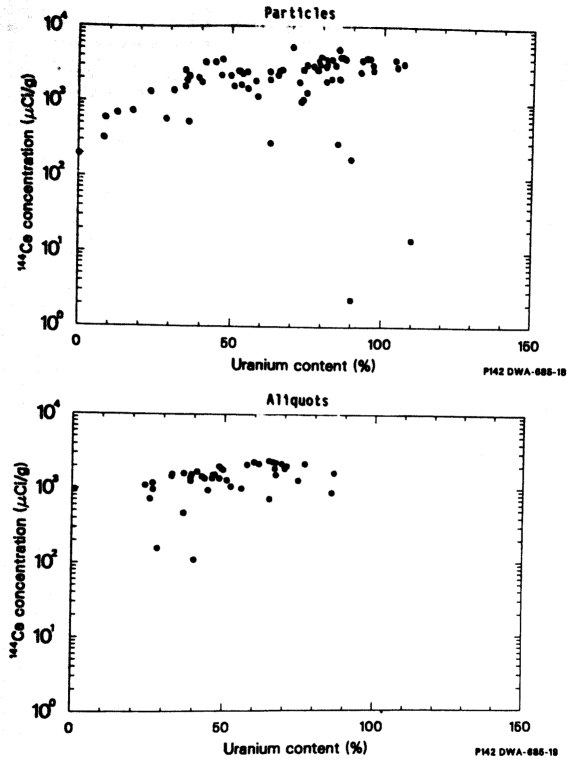


Figure 41. 144Ce/U content correlation for all particles and aliquots analyzed.

Table 29 indicates that $^{106}\mathrm{Ru}$ is retained in the fuel, with only small amounts (\sim 3%) associated with Zr. No general correlation with structural materials (e.g., Ni) was observed, except for particles 9D and 9G which are metallic appearing and contain significant quantities of Ni. A comparison of the $^{106}\mathrm{Ru}$ and $^{\bar{1}25}\mathrm{Sb}$ radionuclide concentrations with the Ni concentration are listed in Table 31. The radionuclide concentrations are 10 to 100 times greater than would be expected for a UO₂ fuel pellet, even at high burnup (6200 MWd/MTU). This behavior appears to be relatively specific to these particles. The retention for ^{106}Ru ranges from 163 to 187 $\mu\text{Ci/mg}$ Ni. For ^{125}Sb , the range is 17 to 20 $\mu\text{Ci/mg}$ Ni. The Ni to ^{106}Ru and Ni to ^{125}Sb atom ratios are 3.8 E+4 and 6.9 E+5, respectively. This indicates that the 106 Ru or 125_{Sb} are located inside the Ni matrix, with many atoms of Ni present for every atom of either radionuclide. Using the retention factors indicated. the core inventory of Ni (~1046 Kg) could retain 1.96 E+5 Ci of 106Ru and 2.09 E+4 Ci of 125 Sb. This is equivalent to 170% of the 106 Ru core inventory and 56% of the 125 Sb core inventory. However, as noted, this is not a general trend and is specific to a few particles. Also it is unlikely that significant amounts of the structural materials would be in contact with the fuel to produce this behavior. Figures 36 and 37 show the correlation between Ni content and 106Ru and 125Sb concentrations.

 $^{125}\mathrm{Sb}$ and $^{90}\mathrm{Sr}$ exhibit significantly different retention characteristics as indicated in Table 29. The $^{125}\mathrm{Sb}$ appears to be present in greater concentrations in the Zr-bearing than in the U-bearing particles, suggesting significant release from the fuel; whereas, $^{90}\mathrm{Sr}$ appears to be principally retained in the fuel. Figure 38 shows the correlation between $^{90}\mathrm{Sr}$ and U content. There is an apparent reduction in concentration at lesser U content. As shown in Table 30, the $^{125}\mathrm{Sb}$ appears to be scavenged by structural material. Figure 39 shows the relationship between $^{125}\mathrm{Sb}$ and U content. No apparent correlation is present. Figure 40 shows the correlation between $^{125}\mathrm{Sb}$ and $^{129}\mathrm{I}$ in order to determine if these radionuclides ($^{129}\mathrm{I}$ and $^{125}\mathrm{Sb}$) are exhibiting similar behavior. No obvious correlation is apparent.

TABLE 37. RETENTION OF TOBRU AND 125Sb BY PARTICLES 9D AND 9G

Concentration	Particle 9D	Particle 9G 94.6
with content (mg/g)	186	
Ag content (mg/g)	16	18
106 _{Ru} content (µC1/g)	4.65 E+4	1.77 E+4
Sample 9 106Ru average concentration ^a (µC1/g)	3.80 E+2	3.80 E+2
Particle/average concentration ratio ^b	1.22 E+2	4.66 E+1
125 _{Sb content}	5.64 E+3	1.61 E+3
Sample 9 average 125Sb concentration ^a (µC1/g)	7.27 E+1	7.27 E+1
Particle/average concentration ratiob	77.6	22.1

a. Average concentration measured for the bulk sample.

b. Ratio of the concentration in the particle to that of the recombined bulk sample.

The Ce and Eu appear to be retained principally in the fuel matrix, as shown in Table 29. However, retention of ^{144}Ce by the Zr material was found on a single particle (Particle 6B). Otherwise, little release from the fuel is indicated. Figure 41 shows the correlations between ^{144}Ce and U content for particles and aliquots. For particles, there is a wide diversity in concentration. Burnup may account for some variation, because the amount of ^{144}Ce produced is directly related to burnup and enrichment. The data indicate a range with relatively high Ce concentrations (~10 $^3\mu\text{Ci/g}$) associated with particles of both high and low U content.

3.4.5 Evidence of Core Material Mixing

The effective enrichment calculations discussed in Section 3.4.3 provide evidence of substantial mixing of core materials (see Table 26). Effective enrichment is calculated by dividing the measured fissile material (235 U) content of each recombined bulk sample by the total U content obtained from the elemental analysis. There were three fuel assembly enrichments in the TMI-2 core. The peripheral fuel assemblies were 2.96% enriched, and the inner fuel assemblies alternated between 1.98% and 2.64% enriched. At the H8 and E9 locations the original enrichments in the fuel assemblies were 2.64% and 1.98%, respectively. The data indicate a variety of enrichments, suggesting significant mixing of the U from the fuel assemblies. The effect is pronounced at E9 where many of the samples indicate the presence of 2.7% enriched fuel material.

A comparison was performed of the measured enrichments for the particles and aliquots from the various samples. A wide range exists, indicating there is no correlation with the original enrichments at the sampled locations (see Table E-11). Chemical and radiochemical evidence of core relocation is further evaluated in Section 4.

3.4.6 Summary of the Cestum Release and Settling Tests

Cesium release, turbidity, and airborne release tests were conducted on 50 g of recombined bulk sample material from Sample 6. The tests were performed on both the as-received core debris materials and on debris which was crushed to alter the particle size distributions and create freshly fractured surfaces. Crushing was intended to simulate the breakup of TMI-2 core material during reactor defueling. Details of these examinations are reported in Reference 30, and the results are summarized below.

- o Crushing the debris increased the concentration of ¹³⁷Cs in solution by a factor of approximately 10, due to the exposure of freshly created surfaces (by crushing) that allowed leaching into the coolant.
- Turbidity examinations indicate that crushing the debris causes little change in the rate of decrease in turbidity. Also, significant reductions in turbidity occur within 24 hours. These data indicate that crushing the debris during defueling will have little impact on turbidity during defueling operations.
- Airborne radioactivity measurements indicate that production of airborne radioactivity during defueling is expected to be significant only when wetted surfaces are near the point of drying out. Minimizing changes in the volume of reactor coolant in tanks or in the core cavity will reduce the exposure of nearly dry radioactive surfaces and consequently the production of airborne radioactivity in the reactor building.

4. CONTRIBUTIONS TO UNDERSTANDING THE TMI-2 ACCIDENT

The influence of the core debris examinations on understanding the TMI-2 accident scenario, the current condition of the core, and radionuclide retention/release from the fuel and core region is discussed in this section. A brief summary of the current understanding of the accident is presented, followed by sections describing (a) the temperature estimates and fuel behavior, (b) evaluation of core relocation, and (c) radionuclide behavior analyses.

4.1 Current Best Estimate of the TMI Accident

The current state of the TMI-2 core and support structure is based on the following examinations and calculations: (a) closed circuit (CCTV) inspections of the upper and lower plenum and core regions; (b) topography measurements of the core cavity; (c) examinations of leadscrews and leadscrew support tubes; (d) mechanical probing of the debris bed to determine the debris depth at 17 locations; (e) measurements from online instrumentation; (f) probing of selected instrument guide tubes with small diameter (piano) wire to determine penetrability of the core from below the reactor vessel; (g) gamma scans of the lower vessel region and reactor coolant system to identify potential fuel deposits outside the active core region; (h) visual examination of the plenum assembly; and (i) computer code calculations and hypotheses.

The known physical condition of the reactor core is summarized in Figure 42. A void is present in the upper region of the core, with a volume of approximately 25% of the total core volume. Becaminations of the microstructure of two control rod drive leadscrews (central and mid-radius) indicate that structural temperatures in the upper plenum ranged from 700 K in the upper regions to 1350 K in the lower regions just above the core. CCTV inspections show nonuniform damage to the bottom of the plenum assembly, ranging from areas with no damage to areas where extensive melting and oxidation of the stainless steel occurred. The damage zones appear to be limited to the lower few centimeters of the plenum assembly.

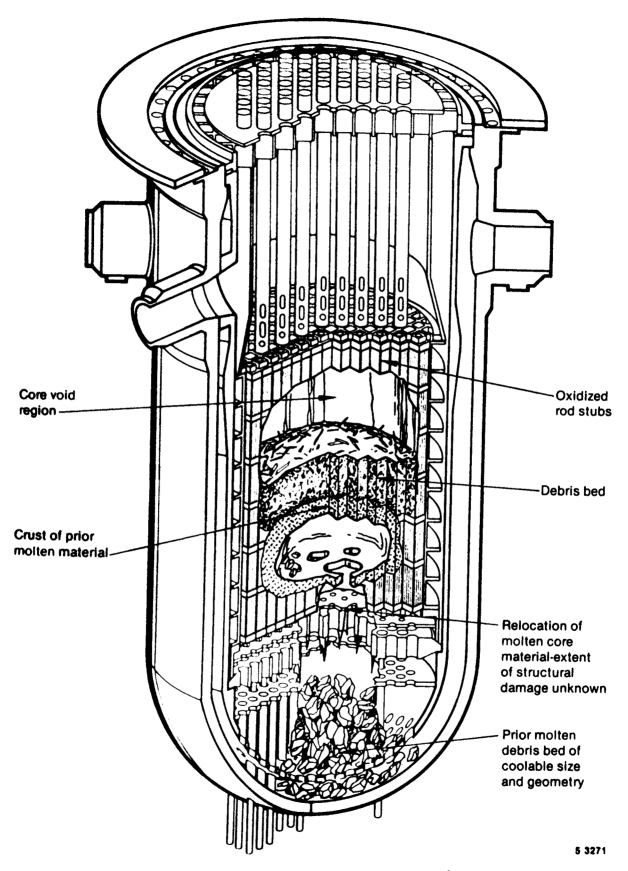


Figure 42. Schematic showing the known and best estimated end-state condition of the TMI-2 core.

A debris bed ranging from 0.8 to 1.0 m thick exists at the bottom of the void. The debris bed is resting on a layer of hard (impenetrable) material at about the mid-core elevation. The condition of the lower core is not known.

CCTV inspections of the lower regions of the reactor vessel indicate that 10 to 20% of the fuel and/or structural materials of the core now reside on the reactor vessel lower head, approximately 2.2 m below the bottom of the original core region. The particle size and texture of the material in the lower plenum appears to vary significantly, ranging from pea-like gravel near the center of the vessel to larger pieces of lava-like material up to about 13 cm in diameter near the periphery. Possible damage to the core support assembly is not known because the central regions of the lower plenum were not visible in the video scans. There is no evidence that the lower reactor vessel head has been damaged.

A scenario for the first four hours of the TMI-2 accident was developed based on (a) end-state conditions of the core and reactor vessel (as described above), (b) measurements of the plant response recorded during the accident, (c) the results of best-estimate calculations of the accident, ³⁴ and (d) severe fuel damage experiments conducted in the Power Burst Facility at INEL. ³⁵ Important events and features of the accident are summarized here for later comparison with the information obtained from examining the core debris grab samples. Details of the accident before core uncovery are discussed elsewhere. ³⁶

Core uncovery began about 100 min after reactor scram. Superheated steam in the hot legs was initially measured at about 113 min. Best-estimate predictions indicate that core temperatures probably increased sufficiently to result in fuel rod ballooning, rupture, and release of gaseous fission products by about 140 min. Radiation measurements indicated a significant release of fission products into the containment at about 142 min. By 150 min, the predictions indicate that peak cladding temperatures rapidly increased due to cladding oxidation and fuel rod temperatures exceeded the melting temperature of zircaloy. The

molten zircaloy would have dissolved a portion of the $\rm UO_2$ fuel pellets. The dissolved fuel and molten zircaloy flowed downward along the fuel rods and eventually froze in lower, cooler regions of the core. The lower level of the previously molten materials was probably coincident with the coolant liquid level which was estimated to be in the lower third of the core. The predictions indicate that a significant portion of the core (1/4 to 1/2) reached temperatures sufficient to melt the cladding and that the core maximum temperature was sufficient to melt the $\rm UO_2$ fuel.

One of the reactor coolant pumps was turned on from about 175 to about 180 min resulting in some coolant flow into the core. Thermal and mechanical shock in the upper regions of the core which consisted primarily of highly oxidized and embrittled rod remenants caused extensive fragmentation and collapse of the rods. The core continued to heat up after the pump transient (as indicated by the in-core thermocouples). The relatively solid mass in the bottom of the core eventually melted through the bottom supporting crust, and molten material flowed into the lower reactor vessel head at about 227 min. A coolable geometry was attained in the lower reactor vessel head, and long term cooling was reestablished at about 16 h into the accident.

Three concepts are unique to this accident scenario. First, based on an understanding of cladding melting and fuel liquefaction, significant portions of the core probably flowed downward into lower portions of the core and solidified within the confines of the lower core boundary prior to the pump transient. This region of liquefied core materials and intact fuel rods could have been of substantial size, from 1 to 2 m in height and from 80 to 90% of the core diameter. Second, because of the thermal properties of the core materials, these large zones containing metallic Zr and ceramic materials probably were not coolable (using reasonable decay heating assumptions), even when surrounded by water. Third, the large solid mass continued to heat up and eventually a significant fraction of the core relocated through the core support assembly into the lower reactor vessel head, where a coolable configuration was attained.

4.2 <u>Impact of Metallurgical Findings on Understanding</u> the Accident Scenario

The metallurgical analysis of 29 particles from the core debris grab samples provided information in several areas that relate to the accident scenario. Estimates were made of core peak temperatures achieved during the accident, some indication of the temperature history was obtained, information on the chemical interactions of the fuel rod material with other core materials on a microscopic scale was obtained, and details of the relocation phenomena were investigated. This section compares results of the metallurgical examinations to the scenario described in Section 4.1. The comparison is presented in two sections: temperature estimates and fuel behavior.

4.2.1 Temperature Estimates

The peak temperature or temperature range achieved during the transient was determined by detailed analysis of the microstructure and elemental composition of the particles. The selection of particles for examination was biased in that particles showing some indication of melting were usually selected. Although most of the 29 particles showed evidence of high temperatures, examination of the bulk samples indicated that a large fraction of the material did not exceed 2000 K for significant lengths of time. Many of the fuel pellet fragments within the melt-bearing particles display little or no equiaxed grain growth, which confirms that the entire core did not experience prolonged times at extremely high temperatures. However, there are particles that show microstructures indicating peak temperatures up to the melting point of the UO, fuel. addition, the evolution of other particles involved oxidation and mixing processes that cannot be readily explained except by lengthy durations under very severe conditions. Both sets of observations are consistent with the postulated accident scenario.

Evidence was found of molten Zr which had flowed down interior cladding surfaces, within the fuel-cladding gap. This phenomenon is

Important in calculating peak temperatures, as well as in estimating fuel liquefaction and fission product release. The longer the molten metallic zircaloy cladding remains in place, the more heat is generated by the exothermic Zr-steam reaction. Containment of molten Zr between UO₂ fuel and an exterior ZrO₂ crust therefore tends to increase peak fuel rod temperatures. Cladding ballooning occurred during the transient, as evidenced by the pieces of deformed cladding found in the debris. The ballooning may have enhanced retention of the molten, unoxidized Zr, thereby driving the temperatures even higher.

4.2.2 Fuel Behavior

Significant interaction (i.e., formation of eutectic mixtures) of the fuel rod materials with structural materials was observed on a microscopic scale, with the eutectic material frequently located in the grain boundaries. Interaction with the Inconel spacer grids seemed to be common, with appreciable amounts of Ni, Fe, and Cr often detected within prior molten regions. This correlates with the CCTV inspection of the upper void, which shows many of the fuel rod assemblies severed at a grid spacer. It is expected that closer inspection will show chemical interaction of the fuel rods with the grid spacer material. Current severe accident analysis computer codes do not adequately model the interaction of the fuel rods with all of the reactor structural components. Nevertheless, these observations on fuel-structural interactions are consistent with results of in-pile and out-of-pile research on severe core damage accidents which are presently being conducted (see Reference 35).

Many particles exhibit distinct prior molten layers of significantly different compositions, where typically one molten mixture solidified (and often oxidized) before being contacted by a second fuel-bearing melt.

Interactions between such layers were found to vary from moderate-temperature diffusion bonding to thermo-chemical dissolution at much higher temperatures. These particles indicate either multiple temperature excursions, separated by oxidation and solidification periods or a prolonged candling sequence during the TMI-2 accident, where droplets of

liquefied fuel had sufficient time to relocate down fuel rod exteriors, chemically react, solidify, and oxidize before other droplets arrived. On certain particles, structural material content differs markedly between layers such that limited lateral movement of the molten material may also have occurred. In any case, these layered particles are not typical of in-pile and out-of-pile experiments that confine fuel liquefaction and solidification to a brief continuous process. Consequently, melt relocation during the high temperature portion of the TMI-2 accident should not be viewed as a single core-wide event, but rather as a complex sequence of interrelated and possibly intermittent phenomena. The short "B" coolant pump operation and the occasional Emergency Core Cooling System flows may have locally rekindled or temporarily accelerated Zr-steam oxidation, thereby creating this particle type.

The occurrence of hyperstoichiometric fuel (independently measured on TMI-2 particles at three laboratories) is another major difference from most in-pile and out-of-pile research efforts to date. These experiments generally involve reducing environments, with Zr oxidation creating strong competition for the oxygen from available steam and from $\rm UO_2$ dissolution. However, hyperstoichiometric Fuel ($\sim \rm UO_{2.5}$) was also discovered by SAS within the Severe Fuel Damage Scoping Test bundle. The fuel oxidation apparently occurred during the SFD-ST cooldown--after zircaloy oxidation and associated hydrogen generation had ceased. This interpretation was aided by a recent thermodynamic calculation that suggests fuel can oxidize beyond $\rm UO_{2.2}$ in high pressure steam ($\sim \rm 100$ atmospheres), but only if little hydrogen is present. He Fuel oxidation in SFD-ST was also tentatively associated with reduced grain boundary adhesion, formation of microcrack networks, and gradual washout of fission products.

4.3 Evaluation of Core Relocation

Knowing the extent of core materials relocation is important to understanding the current condition of the core and postulating a viable scenario for the accident. The metallurgical, chemical, and radiochemical

data provide evidence of extensive mixing, bulk material relocation, and segregation of core materials during the accident. Each of these three types of relocation is evaluated, and their impacts on understanding the condition of the core and accident scenario are addressed.

4.3.1 Mixing of core materials

Evidence of mixing of core materials has been obtained principally from the metallurgical and chemical analyses results (Sections 3.2 and 3.3). The metallurgical data (SEM/EDS and SAS) indicate the presence of structural materials (Cr., Fe, Ni), poison rod materials (Gd, Al), and control rod materials (Ag, In, Cd) in regions of ternary U-Zr-O mixtures, at grain boundaries or on surfaces within the voids of many particles. These data suggest that the structural materials and Al were incorporated into the particles as a result of fuel rod melting and dissolution, or the formation of low melting temperature eutectics.

Elemental analysis of the particles and aliquots indicates the presence of most structural materials in each sample at relatively low concentrations (<1.0 wt%). The uniformity of the structural material concentrations in most particles and aliquots indicates that structural and control rod materials were probably mixed into the debris when the material was molten. These data also suggest that the debris bed was thoroughly mixed for a substantial period of time during the accident to achieve the degree of homogeneity observed for the core constituents.

The distribution of the Gd from the burnable poison rods $(Gd_2O_3-UO_2)$ provides additional evidence of significant mixing of core materials. Only about 13.5 kg of Gd was originally contained in the core, at four widely separated locations (H4, H12, D8, and N8). Gadolinium, however, was measurable in similar concentrations in most of the recombined bulk samples from the H8 and E9 locations. The distances from the H8 and E9 locations to the center of the nearest Gd fuel assembly are 87 cm and 30 cm, respectively. The homogeneity of the Gd concentrations in the samples supports the hypothesis that the core

materials were subjected to significant mixing probably after the accident and after high temperatures had existed because the Gd (either as ${\rm Gd_2O_3}$ or as reduced Gd) was transported to the H8 location and mixed to a concentration similar to what was measured in debris taken from the E9 position. The relocation of the Gd in the core could have occurred via several mechanisms. Gadolinia (melting point ~3000 K) may have been either melted or reduced by molten Zr to Gd and subsequently distributed throughout the debris bed.

Analysis of the mixing of core materials contributes to understanding of core condition as follows:

- o Based on the distribution of core structural materials and their presence in the ternary U-Zr-O structures, the debris bed was at one time probably subjected to significant mixing probably after the high temperature portion of the accident.
- The turbulent mixing of core materials probably occurred during the B-2 pump transient between 175 and 180 min. The relatively uniform concentrations of Gd suggest that the mixing was very vigorous and extensive.

4.3.2 <u>Bulk Relocation of Fuel</u>

Bulk relocation of fuel is suggested from the previous discussion on mixing and from the average enrichment (235 U/total U x 100%) of the recombined bulk samples (see Table 26). The previous discussion concludes that turbulent mixing of core materials probably occurred in order to achieve the observed uniform distribution of core constituents. The 235 U enrichments were also determined for the recombined bulk samples for comparison with the original core enrichments at H8 (2.64 wt%) and E9 (1.98 wt%). The measured enrichments of samples from the H8 location ranged from 2.3 to 2.7 wt% 235 U and at E9 the measured enrichments ranged from 2.5 to 2.8 wt% 235 U. There is no appreciable correlation between the original enrichments, particularly at E9, and measured sample

thoroughly mixed with higher enrichment fuel from other assemblies. For example, the 2.98 wt%-enriched fuel was originally only at the core periphery. Materials with this enrichment are now found in most of the samples examined. The results indicate that significant degradation and mixing of bulk fuel assembly components has occurred.

4.3.3 Segregation of Core Materials

The elements for which segregation or accumulation was indicated, primarily by the chemical analysis results, are Zr, Ag, In, Cd, Gd, and Al. Less than 50% of the anticipated Zr was present in the debris bed at some H8 and E9 locations. Fuel rod temperatures were probably high enough after approximately 150 min for zircaloy cladding to melt, dissolve some UO2, and flow downward along the rods into lower cooler regions of the core. The chemical analysis results support this scenario and suggest that depletion of materials was significant from the upper portion of the fuel rods in the region of the debris bed. Based upon this result, there may be zones lower in the core which contain significant accumulations of the U-Zr-O ternary mixture. The debris bed contains approximately 20% of the core mass and based upon the indicated depletion of Zr, approximately 500 to 2200 kg of Zr with dissolved U may have been deposited near the bottom of the core prior to the B-2 pump transient at 175 min.

The control rod constituents (Ag-In-Cd) are relatively volatile, and based on thermodynamic calculations for conditions similar to TMI, 37 significant amounts of Ag (24%), In (47%), and Cd (100%) could vaporize, condense into aerosols, and be transported out of the core region. The results of the recombined bulk sample analysis indicate the presence of only 7.4 to 10.1% of the initial core concentration of Ag in the core debris grab samples. [Cd and In were not measurable in the recombined bulk samples, probably because the concentrations were below the detection limit.] Evidently, approximately 90% of the Ag has been transported from the debris bed (~400 kg Ag). Less than 3% of the core inventory of Ag left the reactor vessel based upon analyses of components outside the core

[leadscrew surfaces, (see Reference 33) reactor coolant drain tank and reactor building basement, 38] where the Ag would tend to accumulate if transported as a vapor or aerosol out of the core. The silver probably melted and flowed into lower regions of the core or reactor vessel where it probably remains.

The concentration of Ag tends to be higher for the smaller (<1000 μm) particle size fractions possibly because of the increased surface area for condensation on the particles with decreasing particle size. The Ag could also react chemically with the particles and a similar trend would be indicated. The data indicating whether the interaction is physical or chemical do not yet exist. Cd and In, the other principal control rod constituents, were measurable in the particles and aliquots; but in fewer samples than was Ag. These elements exhibit essentially the same behavior as Ag, with higher concentrations associated with the smaller particle size fractions.

Gadolinium, a poison material, was present in most recombined bulk samples. It was distributed relatively evenly, with measured concentrations ranging from 4.7 to 8.2 E-2 wt%. This is surprising as gadolinium constitutes less than 1.1 E-2 wt% of the core and the measured concentrations are significantly higher than the initial core average concentration. No correlation with particle size is apparent. Extrapolation of the measured concentrations to the weight of the debris bed indicates that the core inventory of Gd could be contained in the debris bed. It should be noted that Gd is present only in the middle half (or two-thirds) of the poison rod and would therefore principally have been exposed to accident conditions. It is possible that Gd was concentrated in the debris bed at the sample locations and that large areas with little or no Gd present exist at other locations in the core.

Aluminium, the other primary constituent of the poison rods (excluding 0), appears to have behaved differently than gadolinium. Aluminum was measurable only in the two recombined bulk samples of surface material, at 7 to 20 times the concentration expected if the Al were evenly distributed

the debris bed. Tellurium also was concentrated at the surface the debris bed. In general, the concentration of Al increases with decreasing particle size as indicated in Figure 43. This figure is a comparison between the normalized particle surface area and the concentrations of Al for Samples 1, 3, and 6. The deviations in the general increase in concentration with decreasing particle size may be due to differences between the actual and calculated surface areas of the particles. These data suggest that the transport mechanism may have been vaporization within the high temperature zone and condensation at the top of the cooler rubble bed.

These results contribute to an enhanced understanding of the core condition and accident scenario as follows:

- The distribution of Zr, Ag, Al, and Gd within the debris bed indicates that the unexamined regions in the lower half of the core and lower plenum probably have significantly different compositions than does the debris bed. For instance, most of the Zr and Ag which is missing from the core material in the debris bed is probably in the bottom of the core and lower plenum. The core material in these lower regions is probably depleted in Al and Gd.
- concentration of Al and Te in the surface material of the debris bed suggest that the surface of the debris bed functioned as a trap either during the high temperature portion of the accident or later during natural recirculation.
- o The concentration of Al, Ag, Cd, and In with the smaller particle size fractions suggests that retention was dependent upon particle surface area or that these elements formed particles with the observed size distribution. The most likely mechanisms are probably condensation or chemical reaction of vapors.

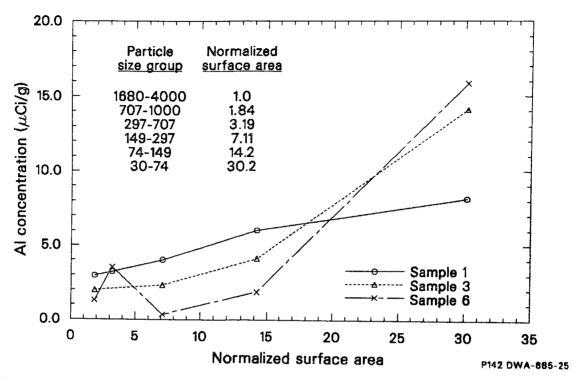


Figure 43. Comparison of Al concentrations with normalized surface area.

4.4 Behavior of Fission Products in the Core

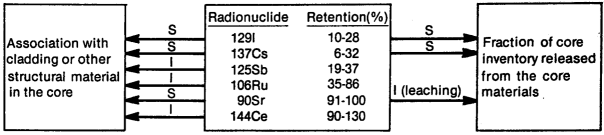
Fission product behavior is defined as the retention/release from the fuel and transport/retention of radionuclides within the reactor pressure vessel and primary cooling system. A block diagram which summarizes the fission product behavior, as determined from analysis of the core debris grab samples, is shown in figure 44. The only measurable radionuclides significantly released from the reactor vessel are \$\frac{137}{25}\$ cs and \$\frac{129}{1}\$. Other radionuclides were retained within the vessel, either in the fuel or associated with the cladding and structural materials. The release of \$\frac{106}{106}\$Ru and \$\frac{125}{125}\$Sb from the fuel is significant. Behavior of each of the radionuclides and their impacts on the current understanding of the core condition and accident scenario is evaluated in this section. The chemistry associated with the observed behavior is evaluated in References 39, 40, and 41.

4.4.1 Cestum and Iodine

(except for the noble gases) and, hence, the most wolatile radionuclides (except for the noble gases) and, hence, the most mobile of the measurable radionuclides. Estimates of radionuclide retention in the core debris, as shown in Figure 44, indicate that the maximum retention in the debris bed for \$^{137}Cs and 129 I is 28 and 32%, respectively. Significant fractions of the core inventory of both radionuclides have been relocated by the reactor coolant to the remainder of the reactor system (see Reference 28). Also, the radiochemical analyses (see Section 3.4) indicate that both radionuclides have become associated with cladding and structural materials at concentrations similar to fuel material (i.e., the structural materials have 129 I and 137 Cs concentrations similar to fuel material).

Comparing the $^{137}\text{Cs/}^{129}\text{I}$ ratios for high fuel content (>50%) samples indicates no correlation between ^{137}Cs and ^{129}I . This suggests that the radionuclides were removed from the fuel material to different extents or subsequent leaching has affected the distribution. Under highly oxidizing conditions as were indicated at TMI by the degree of cladding

Retention in fuel materials



- S Significant relocation
- I Insignificant relocation 5 3266

Figure 44. Schematic showing the behavior (retention/relocation) of various radionuclides inthe TMI-2 core.

oxidation, I^2 would probably preferentially be formed rather than CsI indicating different transport methods. The core debris data tend to support this hypothesis. There appears to be a general increase in ^{129}I associated with the smaller particle size fractions (see Section 3.4); however, no similar relationship exists for ^{137}Cs . If the correlation with the particles size fraction (i.e., surface area) is evidence of vaporization/condensation, these data suggest that ^{129}I vaporized and subsequently condensed on the surfaces of the smaller particles, whereas ^{137}Cs did not deposit as a vapor or was selectively removed by leaching or other mechanisms.

The contributions of the 137 Cs and 129 I results on understanding the core condition and accident scenario are as follows:

- o 134,137Cs and 129I have been significantly removed from the fuel and have become associated with cladding and structural materials or transported outside the vessel, with no correlation between the two.
- o The data suggest that 129 I may have deposited as a condensed vapor on surfaces or reacted with components of the smaller particles size fractions to form an insoluble material not removed by leaching.
- o Previous experiments have indicated that release of Cs and I from the fuel occurs after fuel fracture and quenching. The most significant release probably occurred at 175 min into the accident during the 8-2 pump transient.

4.4.2 Ruthenium

Ru release from the debris was quite significant (14 to 64%). However, $^{106}\rm{Ru}$ is measurable to a minor extent in cladding and structural material samples, suggesting some retention by other core constituents. Significant retention of $^{106}\rm{Ru}$ in the debris was observed for several

isolated particles. For particles 9D and 9G, the Ni to 106 Ru retention was much greater (10 to 100 times) than would be found in a comparable particle of U, indicating that Ni scavenges 106 Ru and incorporates it into the Ni matrix. Extrapolation of the measured retention factors for 106 Ru to the core inventory of Ni indicates the potential retention of the total core inventory of 106 Ru by the Ni present in the core. It is unlikely, however, that all 106 Ru present in the core would be contacted by structural Ni, indicating that this is a less significant potential retention mechanism.

The low retention of 106 Ru in some samples indicates that significant 106 Ru depletion has occurred from the debris bed. No significant amount of this radionuclide has been measured outside the reactor vessel, suggesting that the 106 Ru in the debris bed may have been transported to lower regions in the core. A possible transport mechanism is that the 106 Ru reacted with Zr or structural material and was transported to lower regions of the core with those materials. Although Ru metal has a high boiling point with very low volatity, it has ~10 oxidation states which are highly volatile, unstable chemical compounds (see Reference 40). Formation of those chemical compounds may account for the 106 Ru depletion in the debris bed.

Contributions of the ¹⁰⁶Ru results to understanding the core condition and accident scenario are as follows:

- o Ru is significantly retained in the fuel; however, what has been released probably reacted with structural materials and has been retained within the reactor vessel or near vicinity.
- o lower in the reactor vessel.

4.4.3 Antimony

The 125 Sb results indicate significant release of this radionuclide (63 to 81%) occurred from the debris bed. The release was similar to 137 Cs; however, no significant fraction of the core inventory of 125 Sb has been measured outside the reactor vessel. This radionuclide also behaves similarly to 106 Ru in that it leaves the fuel and associates with cladding and structural materials. The Ni to 125 Sb atom ratio is 125 Sb also is incorporated into the Ni matrix. Extrapolation of the measured retention factors of Ni for 125 Sb indicates that as much as 56% of the core inventory of 125 Sb could have been retained by structural materials containing Ni.

The data suggest that the missing portions of the core inventory may have reacted with Zr or structural material and have been transported to lower regions of the core when these materials were relocated.

The contribution of the \$125\text{Sb}\$ results to understanding the core condition and accident scenario is that a large fraction of the \$125\text{Sb}\$ inventory of the debris bed (60 to 80%) has not been accounted for. The data indicate that the \$125\text{Sb}\$ reacted significantly with structural materials and may have been deposited on those materials within the core region or has been transported to lower regions in the reactor vessel with the relocated cladding and structural materials.

4.4.4 Strontium

 90 Sr has been significantly (> 90%) retained in the fuel and has not become associated with cladding or structural materials, as determined from the analyses in Section 3.4. However, a continuous addition of 90 Sr to the reactor coolant has been measured (see Reference 26), suggesting that this material leaches directly into solution and does not react with Zr or cladding significantly. No significant accumulations of 90 Sr have been measured outside the core. The oxidizing conditions at TMI would tend

to form SrO (see Reference 40), which has very low volatity. Formation of this compound with no subsequent reduction with Sr may be the reason for the low volatility of the 90 Sr at TMI.

The contribution of the 90 Sr results to understanding the core condition and accident scenario is that 90 Sr was retained in the fuel and was not released during the accident. The data suggest that 90 Sr is not an immediate hazard during a severe core damage accident similar to TMI-2 and is only slowly leached from the fuel into the coolant.

4.4.5 Cerium

 $^{144}\mathrm{Ce}$ appears to have been completely retained (90 to 130%) in the reactor vessel. The results indicating retentions of greater than 100% may be due to localized variations in burnup or possible relocation of $^{144}\mathrm{Ce}$ from the fuel. An evaluation of $^{144}\mathrm{Ce}$ relocation was performed in Section 3.4, indicating that $^{144}\mathrm{Ce}$ was released from the fuel and has become associated with cladding and structural materials in the core. $^{144}\mathrm{Ce}$ probably formed CeO, a relatively high volatity compound (see Reference 37), from $\mathrm{Ce}_2\mathrm{O}_3$. CeO would probably be produced by the reduction of $\mathrm{Ce}_2\mathrm{O}_3$, a very stable, low volatility compound. The oxidation of the cladding and core materials would indicate there was a generally oxidizing environment; however, the $^{144}\mathrm{Ce}$ data suggest the presence of a time during the accident when a reducing atmosphere was present in the debris bed to produce the observed $^{144}\mathrm{Ce}$ mobility.

Contributions of the ¹⁴⁴Ce results to understanding the core condition and accident scenario are that no significant release of ¹⁴⁴Ce from the core occurred. What little did escape from the fuel matrix reacted with nearby cladding and structural material and was retained in the reactor vessel. The higher concentrations of ¹⁴⁴Ce in the debris bed may indicate the presence of periods of time during the accident when a reducing atmosphere was present.

5. OBSERVATIONS/CONCLUSIONS

Principal observations and conclusions that have been obtained from examining the core debris grab samples include results of the physical, metallurgical, chemical, and radiochemical analyses and correlation of these results with current information on the known core condition and accident scenario. The principal observations/conclusions based on a total sample size of 0.001% of original core inventory include the following:

- o Approximately 80 wt% of the debris material is larger than 1000 µm. The largest particles were up to 20 mm.
- o five types of particles were identified based on visual appearance: (a) apparent fuel pieces; (b) cladding chunks; (c) foamy/porous, prior molten material; (d) particles that are a composite of fuel and prior molten material; and (e) metallic appearing, prior molten particles.
- Many of the particles examined contain regions of prior molten $(U-Zr)O_2$, indicating peak temperatures greater than 2800 K. Also, there are a few examples of prior molten material that are almost pure UO_2 , indicating temperatures near 3100 K.
- o Much of the fuel examined was essentially unrestructured indicating that peak temperatures did not exceed 2000 K for any significant length of time.
- o Several independently analyzed fuel-bearing particles contain oxygen concentrations in excess of 70 at.%. Such extensive fuel oxidation probably occurred during steam cooling with high steam pressures and very little hydrogen generation from metal oxidation.
- o Regions of prior molten U-Zr-O usually contain at least some trace of non-fuel rod material (Al, Cr, Fe, Ni). These elements

often were found at grain boundaries or in voids. This indicates significant interaction between fuel rods and grid spacers, and possibly other stainless steel components.

- o Control rod materials (Ag, In, Cd) were not as commonly found as structural materials. Ag apparently had less of a tendency to mix or interact with fuel rod materials than did the structural materials.
- The presence of layered, prior molten particles indicates either a prolonged candling sequence or multiple temperature ramps occurred during the fuel liquefaction/relocation period of the accident.
- o Evidence that molten metallic U and Zr flowed inside ballooned cladding was observed, as well as candling on the cladding outside surface.
- o Significant Zr depletion (\leq 50%) has occurred in the majority of the debris samples analyzed, indicating significant relocation of Zr from the rubble bed.
- o The majority of particles and aliquots examined contained a mixture of most core constituents, indicating significant mixing and disruption of the original core. This may have occurred either during the high temperature portion of the accident or later during natural recirculation.
- o Concentrations of Ag-In-Cd (volatile core materials) generally show increases for the smaller particle size fractions and suggest that a retention mechanism associated with surface area may have been operable for these materials.
- o Significant depletion of Ag (~90%) from the core debris bed has occurred, suggesting possible transport to regions lower in the reactor vessel.

- o Wide distribution of the poison material, 'Gd, indicates that significant relocation of core materials has occurred.
- Al and le are concentrated at the surface of the debris bed, suggesting that this portion of the bed may have functioned as a retention zone or trap for some material either during the high temperature portion of the accident or later during natural recirculation.
- o Structural materials are well mixed in the debris bed at concentrations 30 to 50% of their original concentrations in the fueled region of a fuel assembly. The data indicate that both significant mixing of core constituents and little contamination of the debris with other structural components (end boxes, etc.) occurred.
- o Significant amounts (70 to 80%) of all $^{137}\mathrm{Cs}$ and $^{129}\mathrm{I}$ has relocated from the debris bed.
- o The 106Ru, 125Sb, 137Cs, and 144Ce at the E9 core location generally are at higher concentrations than at H8. The 137Cs concentrations are 18 to 85% higher at E9.
- o Portions of 106 Ru have left the fuel and have been retained in other materials (i.e., structural materials).
- o 125 Sb has been significantly released from the fuel and has associated with non-fuel debris.
- o 106 Ru and 125 Sb appear to have associated with Ni-containing materials, suggesting a retention mechanism for these radionuclides.
- o Sr appears to be almost completely retained in the fuel, with some leaching into the reactor coolant.

- Radionuclides were released from the fuel in the approximate order of volatility of the elemental constituents, except for $^{106}_{\rm Ru}$ and $^{90}_{\rm Sr}$. $^{106}_{\rm Ru}$ release is higher and $^{90}_{\rm Sr}$ release is lower than expected, probably due to the formation of higher or lower volatile oxide forms.
- o 144 Ce concentrations in the samples are higher than expected, based on the average core burnup. Some particles with little or no U content contain 144 Ce. These data suggest relocation of 144 Ce in the debris bed, possibly by formation of a volatile oxide.
- o 129I content in the debris tends to correlate with surface area in the small particle size fractions and, therefore, may suggest a surface area dependent retention mechanism.
- o Evidence of significant turbulent mixing of the liquefied core material has been observed.
- o Segregation of Zr, Al, Ag, and Gd has been observed, suggesting that lower areas of the core may have different compositions than observed in the debris bed.
- o Ru and 125 Sb may have been transported to lower in the core and associated with liquefied Zr and structural material.
- o There is no correlation between the enrichment of materials examined and the original (H8 and E9) fuel assembly enrichments, indicating significant physical relocation of the bulk core materials (fuel assemblies) has occurred.

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